Crude oil is rich in hydrocarbons, mainly paraffins (alkanes), naphthenes (cyclo-alkanes) and aromatics, with no or very little olefins (alkenes) present. Compounds of the same skeletal structure, but containing heteroatoms like sulphur, nitrogen, oxygen and metals like nickel and vanadium, are also present. Sulphur (≤ 5%) and nitrogen (≤ 0.5%) are the heteroatoms that affect refining the most. In addition to these, corrosion problems may result from oxygenates in the form of carboxylic acids (≤ 3 mg KOH/g) and salts in the form of MgCl₂ and CaCl₂ (≤ 350 μg·g⁻¹). The importance of crude density and distillation characteristics on refining has also been highlighted.

1. Introduction

Crude oil is a collective term used to describe a hydrocarbon rich mixture of compounds that is usually found as a subterranean deposit that accumulated over millions of years. The physical and chemical characteristics of crude oil vary widely from one production field to another and even within the same field. Colloquially terms like “light”, “heavy”, “naphthenic”, “paraffinic”, “sweet” and “sour” are used to characterise crude oil in terms of its boiling range, composition and sulphur content. These are important properties, because they have a big impact on the refinery, but it belies the true complexity and diversity of crude oil. The crude oil composition determines its price and although specific oil types, like Brent crude, are used as an international measure of oil price, more difficult to refine oils command a much lower price. The selection and scheduling of crude oil feed to a refinery is a highly specialised field in its own right, and it directly affects the production economics.

It is important to discuss crude oil properties, because refining technology has been developed in response to the differences between crude oil properties and final product specifications. Furthermore, Fischer-Tropsch derived syncrude is different from crude oil in many respects and it is important to understand these differences, because they relate to the

---

a The terms “sweet” and “sour” originated as measure of corrosiveness. Crudes with more than 6 μg·g⁻¹ dissolved H₂S were considered “sour”, because it led to observable storage tank corrosion. These terms are no longer applicable and it is strictly speaking incorrect to directly relate “sweet” and “sour” to sulphur content.
ability of refining technologies to deal with Fischer-Tropsch products. Crude oil will be discussed in terms of its composition and characterisation in sufficient detail to enable these objectives to be met.

2. Crude oil composition

The compounds and compound classes present in the lighter fraction of crude, which is typically the fraction that can be recovered by atmospheric distillation, can be identified by chromatographic and spectroscopic techniques. The heavier fraction is more difficult to characterise, because it contains molecular structures with molecular masses that can exceed 10 000 g·mol⁻¹. The heavy fraction is classified based on solubility (Figure 1),² with *maltenes* being soluble in *n*-heptane and *asphaltenes* being soluble in benzene.³ The usefulness of this method of classification is doubtful, since it is unlikely that the chemical nature of the molecules is adequately captured. The point has aptly been made by Gray⁴ that these heavier fractions are better described as heteroatom species with limited hydrocarbon character, which tend to associate (almost like Velcro™) making separation by solubility and precipitation untenable. The popular notion of heavy crude oil molecules being polycondensed aromatic hydrocarbons (almost like graphite) does not explain the properties and conversion propensity of heavy crude.

![Figure 1. Classification procedure for heavy crude oil fractions (> 360°C boiling fraction).](image)

² The asphaltene content is determined by precipitating the asphaltenes from the benzene solution with a light paraffin. Industrially propane or butane is used, but practice varies and in the USA *n*-pentane is used. This can lead to differences in the reported asphaltene content and composition of a crude oil. If a light paraffin is used, some heavier paraffins may co-precipitate with the asphaltene fraction, reducing its reported aromatic content.
The asphaltene content of crude oil varies from 0.1% to more than 20% depending on the production field. Asphaltenes contain high concentrations of heteroatoms, like sulphur and nitrogen, as well as metals, like nickel and vanadium. Due to the heavy nature of the asphaltenes, their characterisation is mostly done by techniques used in polymer analysis, like size exclusion chromatography. In refineries equipped with a deasphalting unit, the asphaltenes are recovered as asphalt and not upgraded to transportation fuel.

Crude oil can be classified by chemical composition, density, viscosity and distillation characteristics to name a few. The classification system based on composition refers to only the hydrocarbon nature of oil, namely paraffinic, naphthenic or mixed depending on the compound class that is dominant. The other classification systems find utility in giving an indication of the straight run content of the various distillation ranges, which are important in determining the product slate that can be expected. This might have been useful in the past, but it is doubtful whether these classification systems have adequate value for present refineries, where the true composition of the crude is more important in determining the refining pathways. The point is aptly made by Speight: “Clearly the use of one physical parameter, be it API gravity or any other physical property for that matter, is inadequate to the task of classifying conventional petroleum, heavy oil, and tar sand bitumen.”

2.1. Hydrocarbons

The main hydrocarbon compound classes present are paraffins (alkanes), naphthenes (cycloalkanes) and aromatics, with very little or no olefins (alkenes) present. The presence of dienes and alkynes are extremely unlikely. The paraffins found in crude oil tend to have a high linear hydrocarbon content. This has implications for the fuel quality of straight run products. Light straight run (LSR) naphtha has a low octane number and can generally not be included in motor-gasoline without further refining. Conversely, straight run middle distillates, especially from paraffinic crudes, only needs to be desulphurised (sweetened) to yield good jet fuel or high cetane diesel.

The naphthenes most frequently encountered in crude oil have 5- or 6-membered rings and multiple rings may be present. In some cases systems with 4- and 5-membered
rings are found, which serve as biochemical markers. The naphtha range naphthenes make good reforming feedstock, while middle distillate range naphthenes make good jet fuel.

Mono-, di- and polynuclear aromatics are found in high concentrations in most crude oils. In the motor-gasoline range, these molecules contribute significantly to the octane number of straight run products and the gravimetric energy density of jet fuel. However, the higher homologues are not desirable.

### 2.2. Sulphur containing compounds

Sulphur is the heteroatom most commonly found in crude oil and the total sulphur is usually in the range 0.1% to 5% depending on the crude and seldom falls outside the range 0.05-6.0%. It is also the least desirable heteroatom from a transportation fuel perspective. The distribution of sulphur compounds is not equal over the distillation range and the sulphur content generally increases with boiling point (Table 1).

<table>
<thead>
<tr>
<th>Distillation range (°C)</th>
<th>Sulphur content (%)</th>
<th>Thiols</th>
<th>Sulphides</th>
<th>Thiophenes</th>
<th>Other †</th>
</tr>
</thead>
<tbody>
<tr>
<td>70-180</td>
<td>0.02</td>
<td>50</td>
<td>50</td>
<td>trace</td>
<td>-</td>
</tr>
<tr>
<td>160-240</td>
<td>0.2</td>
<td>25</td>
<td>25</td>
<td>35</td>
<td>15</td>
</tr>
<tr>
<td>230-350</td>
<td>0.9</td>
<td>15</td>
<td>15</td>
<td>35</td>
<td>35</td>
</tr>
<tr>
<td>350-550</td>
<td>1.8</td>
<td>5</td>
<td>5</td>
<td>30</td>
<td>60</td>
</tr>
<tr>
<td>&gt;550</td>
<td>2.9</td>
<td>trace</td>
<td>trace</td>
<td>10</td>
<td>90</td>
</tr>
</tbody>
</table>

† = Benzothiophenes and heavy sulphides.

The sulphur can be present in an inorganic form as elemental sulphur, carbonyl sulphide (COS) or hydrogen sulphide (H₂S), or it can be present in an organic form. The organic classes most often encountered are sulphides, thiols (mercaptans) and thiophenes.

---

6 Hydrocarbons with 4-membered rings give an indication of the crude oil origin, since such compounds are remnants of the structure of the living matter from which it has been derived.

7 The sulphur content of the crude is related to its origin. For example, on a water-free basis plants contain 0.1-0.4%, molluscs about 0.4% and marine algae percentage levels. There are oil sources with a very high sulphur content, like 14% sulphur crude from Rozel Point in Utah, USA, where it is not clear whether the sulphur only comes from organic material.

8 Free sulphur is not often present in crude oil, but it can be present as a suspension or it can be dissolved in the oil. H₂S is found in the reservoir gas, but the amount dissolved in the oil is typically less than 50 μg g⁻¹.
Thiols have the general formula R-SH and are sulphur equivalents of alcohols. Due to the presence of the S-H functional group the thiols are acidic in behaviour (much more than alcohols). They are also extremely malodorous. The thiols are mainly present in the lighter boiling fractions, but their concentration in crude oil is generally quite low.

Sulphides can either be mono-sulphides of the general formula R-S-R′, or disulphides of the general formula R-S-S-R′. Both acyclic and cyclic sulphides are present, including sulphides that are attached to aromatic rings, like thiaindane (Figure 2).

Figure 2. Molecular structure of thiaindane.

Thiophenes are aromatic compounds containing sulphur as part of the ring structure and may take the form of thiophene, benzothiophene, dibenzothiophene and even heavier thiophene derivatives (Figure 3). Most of the sulphur in crude oil occurs in this form. The thiophenes and especially the sterically hindered dibenzothiophenes present a significant challenge to deep hydrodesulphurisation (HDS) to reduce the sulphur content to less than 10 μg·g⁻¹ for transportation fuel. This led to some other avenues being explored in order to deal with these difficult to HDS compounds.

Figure 3. Molecular structures of thiophene derivatives.

2.3. Oxygen containing compounds

The oxygen containing compounds in crude oil is probable the least discussed of all the heteroatom compounds. This is understandable, because oxygenates are generally present in lower concentrations (0.05-1.5%) than sulphur compounds and apart from carboxylic acids that are present in some crudes, the other oxygenates have little impact on transportation fuel quality or the refining processes. Most of the oxygenates are converted to hydrocarbons during hydrotreating, but information on hydrodeoxygenation (HDO) specifically is quite
limited.\textsuperscript{(13)} The main oxygenate classes present in crude oil are furanes, phenols, esters and carboxylic acids.\textsuperscript{h}

Furans are the oxygen equivalent of thiophenes and contains oxygen as part of an aromatic ring structure (as Figure 3, but with O instead of S).

Phenols are aromatic compounds of the general formula Ph-OH. Although phenol itself is responsible for transportation fuel elastomer incompatibility, the alkyl derivatives of phenol are used as oxidation inhibitors\textsuperscript{(14)(15)(16)} in fuel and quite useful at the concentrations normally present in crude oil.

Esters have the general formula R(CO)OR’ and are considered quite benign in transportation fuel.\textsuperscript{i} Both aliphatic and aromatic esters can be found in crude.

Carboxylic acids can cause corrosion problems when processing crude at high temperature.\textsuperscript{j} The acids are not evenly distributed over the whole boiling range, but have a peak concentration in the 400-450°C range.\textsuperscript{(6)} In lighter cuts some linear aliphatic acids can be found, but the acids are mostly naphthenic acids (5- and 6-membered ring cycloalkane carboxylic acid derivatives) and can include a variety of polycyclic naphthenic acids (Figure 4).\textsuperscript{(17)} The acid number of crudes typically varies from 0 to 3 mg KOH/g.

\textbf{Figure 4. Some of the naphthenic acids identified in Bohai crude oil.}

Although naphthenic acids appear to be the most prevalent type of carboxylic acid, some crude oil deposits are rich in aliphatic carboxylic acids (>0.1%), such as the crude oil from the Kutei Basin in Indonesia.\textsuperscript{(18)}

\textsuperscript{h} Amides are also present, which contains the -(C=O)-N< function group. The amides will be discussed with the nitrogen compounds.

\textsuperscript{i} The methyl esters of fatty acids in various plant derived oils form the main constituent of biodiesel.

\textsuperscript{j} Although crude oil generally does not contain some of the very aggressive short chain aliphatic acids, the heavy naphthenic acids that are present in the crude become quite aggressive at temperatures of 350-400°C, which are often encountered during desalting and lubricating oil production. At such temperatures carbon steel is readily corroded and special alloys are required to deal with acidic crudes.
Crude can also contain water, which can be seen as a form of inorganic oxygen. The water is found partly as dissolved water and partly as emulsions. The water content can range from traces to about 2% depending on the crude.

2.4. Nitrogen containing compounds

Nitrogen containing compounds are mostly found in crude oil fractions boiling above 250°C. The nitrogen content depends on the crude origin, but is generally less than 0.5%, although it may be as high as 2%. Basic, neutral and acidic nitrogen compounds are present, with the main nitrogen containing compound classes being amines, amides, pyrroles and pyridines. The basic nitrogen containing compounds reduce the acidity of refining catalysts and are especially undesirable. Nitrogen compounds may also contribute to gum formation in products such as domestic fuel oil.

Amines are basic nitrogen compounds and are found as primary (R-NH₂), secondary (R-NH-R′) and tertiary (R-NR′R″) amines.

Amides are acidic nitrogen compounds containing the –(C=O)-N functional group and are found as both aliphatic and aromatic compounds.

Pyrroles are aromatic compounds containing nitrogen as part of a 5-membered ring structure and may be present as pyrrole, indole, carbazole and even heavier derivatives of pyrrole (Figure 5). These are all neutral compounds, despite the presence of the N-H functionality.

---

Figure 5. Molecular structures of pyrrole derivatives.

---

k Such emulsions may be caused by the soaps of the carboxylic acids present in the crude oil, which are known to be surface-active compounds.
l Crude oil also contains nitrogen compounds of the porphyrine family, which chelates some metal ions and will be discussed with the metal containing compounds.
m This is the result of the lone-pair electrons on nitrogen being delocalised in the aromatic π-cloud and they are consequently not available for sharing with acids. In most of the resonance structures of pyrrole, the nitrogen carries a positive charge.
Pyridine and its derivatives like quinoline, isoquinoline and acridine (Figure 6) are basic nitrogen compounds, with the nitrogen forming part of a 6-membered aromatic ring. These compounds and their heavier derivatives occur throughout the distillation range.

Figure 6. Molecular structures of pyridine derivatives.

2.5. Metal containing compounds

The main metal impurities in crude oil are nickel, vanadium and to a lesser extent iron too.\(^{(7)}\) These metals are generally found in the heaviest fractions and end up as deposits in the residue processing units of a refinery. Some of the metals are trapped in molecules of the porphyrine family (Figure 7), which act as efficient chelating agents for ions like Ni\(^{2+}\) and VO\(^{+}\).\(^{(2)}\) The metal content is very crude dependent and may vary from less than 1 μg·g\(^{-1}\) to more than 1000 μg·g\(^{-1}\). However, it should be noted that not all Ni and V are contained in porphyrine structures.\(^{(19)}\)

Figure 7. Porphyrine structure for chelating metal ions (M) like nickel and vanadium.

Metals are also introduced in the form of salts of sodium, magnesium and calcium, usually as the chlorides (NaCl, MgCl\(_2\) and CaCl\(_2\)), but calcium can also be present as the sulphate or carbonate (CaSO\(_4\) and CaCO\(_3\)).\(^{(6)}\) The salt levels vary, with most crudes having a salt level of between 25-350 μg·g\(^{-1}\) of NaCl equivalent. The major portion of salt is found in the residues, causing problems in the refinery, like fouling of burners and deterioration of product quality. Furthermore, magnesium and calcium chlorides start to hydrolyse at 120°C (Equation 1), a reaction that proceeds quite readily at higher temperatures and that can lead to severe corrosion in the presence of water.
MgCl₂ + 2 H₂O → Mg(OH)₂ + 2 HCl … (1)

There are other metal impurities too, which can be present at low concentrations, like mercury and arsenic. Impurities like Si, Cu, Pb and P may also be introduced during refining and recycling of petroleum wastes.⁷ Organometallic soaps of Zn, Ti, Ca and Mg may also be present as surface-active compounds in the oil-water interface.⁴

3. Crude oil physical properties

3.1. Density

The density of crude oil is reservoir dependent and generally falls between 800 and 1000 kg·m⁻³. It gives an indirect indication of the crude composition, giving rise to the classification of crude based on density (Table 2),⁶ which affects its price. In the oil industry the density is often expressed in degrees API and the conversion to SI units is given by Equation 2, where density (ρ) is in units of kg·m⁻³:

°API = 141.5 / (0.001·ρ) – 131.5 … (2)

<table>
<thead>
<tr>
<th>Classification</th>
<th>Density range (kg·m⁻³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Light crudes</td>
<td>&lt; 825</td>
</tr>
<tr>
<td>Medium crudes</td>
<td>825-875</td>
</tr>
<tr>
<td>Heavy crudes</td>
<td>875-1000</td>
</tr>
<tr>
<td>Extra-heavy crudes</td>
<td>&gt; 1000</td>
</tr>
</tbody>
</table>

3.2. Pour point

The pour point of crude oil is important from a processing point of view. It essentially defines the temperature that is required to pump the crude.⁵ Since crude oil is a complex mixture of compounds, it does not congeal at a specific temperature, but different compounds
crystallise out of solution at different temperatures. The thermal history of crude also plays a role in determining the pour point and must be known. Heating a crude to 45-65°C redissolves some of the paraffinic seed crystals, thereby lowering the pour point, while heating it to 100°C would again raise the pour point due to loss of light material by evaporation.

In practice pour point is determined after preheating to 45-48°C, as defined by the ASTM D97(20) method. Crude oil pour points usually range between –60 and +30°C.\(^6\)

### 3.3. Viscosity

The viscosity of crude mainly influences pumping cost, because it determines the pressure drop in pipelines and refinery units. The viscosity-temperature relationship depends on the crude composition, with paraffinic crudes showing a rapid increase in viscosity with a decrease in temperature, while naphthenic crude have a more gradual response. The viscosity of crude oils varies over a wide range, having values of less than 10 to more than 5000 cSt.\(^o\)

### 3.4. Vapour pressure

The vapour pressure of crude oil as it is produced can be as high as 2 MPa. Storage and transportation of crude with such a high vapour pressure is uneconomical. To reduce the vapour pressure, the crude oil goes through a number of pressure reduction stages, which are basically flash drums in series. The pressure is reduced from the wellhead pressure to less than 0.1 MPa. The associated gas\(^p\) obtained during the pressure reduction is separated from the crude, leaving the crude oil with a vapour pressure of less than 0.1 MPa.

### 3.5. Distillation

Of all the physical properties, the distillation curve probably has the biggest impact on refining, since it determines the distribution of straight run\(^q\) products. The actual distillation

---

\(^a\) This is an oversimplification, since “pumpability” is affected by pumping itself, which increases fluidity by inhibiting crystallisation processes.

\(^b\) Viscosity measurements refer to the kinematic viscosity at 20°C in centistokes (mm²·s\(^{-1}\)), as measured by the ASTM D445 method.

\(^p\) The production of associated gas is unavoidable and recovery is not always economical, despite it sometimes being a significant fraction of the product. For example, the reservoirs in the Middle East have a high associated gas production, typically in the order of 0.14 kg gas per kg of crude. That is why it makes economic sense to construct liquefied natural gas (LNG) and gas-to-liquids (GTL) plants there.

\(^q\) A straight run product is a product obtained by distillation only.
fractions (cuts) that will be produced on the refinery are determined by the refinery design, as well as the specifications of the fuel types to be produced. Typical cuts are given in Table 3. (6)

Table 3. Typical distillation cuts produced in a refinery.

<table>
<thead>
<tr>
<th>Description</th>
<th>Hydrocarbon range</th>
<th>Distillation range (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas</td>
<td>C₃-C₄</td>
<td>&lt; 20</td>
</tr>
<tr>
<td>Light gasoline</td>
<td>C₅-C₆</td>
<td>20-80</td>
</tr>
<tr>
<td>Heavy gasoline</td>
<td>C₇-C₁₀</td>
<td>80-180</td>
</tr>
<tr>
<td>Kerosene</td>
<td>C₁₀-C₁₄</td>
<td>160-260</td>
</tr>
<tr>
<td>Gas oil</td>
<td>C₁₅-C₂₂</td>
<td>260-360</td>
</tr>
<tr>
<td>Residue</td>
<td>C₂₂⁺</td>
<td>&gt; 360</td>
</tr>
</tbody>
</table>

Table 4. Distillation characteristics of Arabian Light crude oil, as example to show its relationship to refining requirements.

<table>
<thead>
<tr>
<th>Property</th>
<th>Gas</th>
<th>Light gasoline</th>
<th>Heavy gasoline</th>
<th>Kerosene</th>
<th>Gas oil</th>
<th>Residue</th>
</tr>
</thead>
<tbody>
<tr>
<td>Distillation range (°C)</td>
<td>&lt; 20</td>
<td>20-80</td>
<td>80-180</td>
<td>180-250</td>
<td>250-370</td>
<td>&gt; 370</td>
</tr>
<tr>
<td>Fraction (vol %)</td>
<td>1.7</td>
<td>6.3</td>
<td>17</td>
<td>11.7</td>
<td>21.9</td>
<td>41.4</td>
</tr>
<tr>
<td>(mass %)</td>
<td>1.1</td>
<td>4.9</td>
<td>14.8</td>
<td>11</td>
<td>21.9</td>
<td>46.3</td>
</tr>
<tr>
<td>Density @ 15°C (kg·m⁻³)</td>
<td>-</td>
<td>659</td>
<td>747</td>
<td>798</td>
<td>854</td>
<td>956</td>
</tr>
<tr>
<td>Sulphur content (mass %)</td>
<td>-</td>
<td>0.024</td>
<td>0.036</td>
<td>0.16</td>
<td>1.4</td>
<td>3.2</td>
</tr>
<tr>
<td>Aromatic content (mass %)</td>
<td>-</td>
<td>1.5</td>
<td>14.3</td>
<td>20.7</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>RON</td>
<td>-</td>
<td>60.6</td>
<td>23.5</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Pour point (°C)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-48</td>
<td>-6</td>
<td>10</td>
</tr>
</tbody>
</table>

Knowing the volume fractions going to the various distillation columns and refinery units if a specific crude is to be processed, enables proper scheduling of the crude. (7) It also allows an economic analysis of intended crude purchases, by matching the crude oil

---

(6) Every refinery unit has a production capacity range, the upper limit set by the maximum operating capacity of the design and the lower limit set by the turndown ratio of the design. The crude oil composition of the feed to the refinery must be such that all refinery units can be operated within their design limitations. The economics of refinery operation is determined by selecting the crude slate in such a way that all units can be operated as close as possible to their maximum capacity given the final product market constraints.
characteristics to the market requirements for final products. The cost of processing the 
crude oil into transportation fuels has to be factored into these calculations. The straight run 
product yield gives an indication of the amount of work that has to be done by the conversion 
units in the refinery to produce marketable products (Table 4). This topic will be explored 
in much more detail when dealing with crude refinery design in the next chapter.

4. Literature cited

(12) Takahashi, A.; Yang, F. H.; Yang, R. T. New sorbents for desulfurization by \( \pi \)-


(14) Rosenwald, R. H.; Hoatson, J. R.; Chenicek, J. A. Alkyl phenols as antioxidants. *Ind.

(15) Wasson, J. I.; Smith, W. M. Effect of alkyl substitution on antioxidant properties of

(16) Nixon, A. C.; Minor, H. B.; Calhoun, G. M. Effect of alkyl phenols on storage and

(17) Qi, B.; Fei, X.; Wang, S.; Chen, L. Study on distribution and composition of carboxylic

(18) Gallup, D. L.; Curiale, J. A.; Smith, P. C. Characterization of sodium emulsion soaps

(19) Dunning, H. N.; Moore, J. W.; Bieber, H.; Williams, R. B. Porphyrin, nickel, vanadium,

(20) ASTM D97. *Standard test method for pour point of petroleum products.*
CHAPTER IV
Crude Oil Refineries

The history of crude oil refining is used to illustrate how refinery design is influenced by raw material cost and availability, as well as product specifications and market demand. Four generations of refinery design are described: Distillation only (1st generation), topping-reforming (2nd generation), topping-reforming-cracking with vacuum distillation and optionally visbreaking (3rd generation), and topping-reforming-cracking-visbreaking-alkylation-isomerisation with vacuum distillation (4th generation). The change drivers, namely raw materials, profitability and market demands, are resulting in certain trends that will determine how future crude oil refineries will look. These trends are: Intensification of residue upgrading, making more hydrogen available, increasing refinery complexity, addition of renewable fuels and the possibility to produce synthetic fuels. As consequence, future crude oil refineries will process more olefin intermediates and be more hydrogen rich.

1. Introduction

The purpose of a refinery is to transform a raw material into a more valuable end product that meets market demands. This is a general definition and holds true for all refineries.\(^a\) Since the production of crude oil started in the 1850’s, the market for crude oil derived products changed dramatically. As a natural consequence, crude oil refineries had to change in design and complexity to keep up with the changes in the market. As crude became available from more and more sources, refinery design not only had to cope with changes in products, but also with changes in feed.

In the middle of the nineteenth century a market was created for kerosene (lamp oil), when it was realised that it could be used in wick lamps, because it burns with a bright smokeless flame without leaving a residue.\(^{b,1}\) The first crude oil refineries consisted of a

\(^a\) The combination of processes to transform metal bearing ores into metals is also rightly called a refinery.

\(^b\) It is wrong to imply that kerosene was initially made from crude oil, it was not, it was made from coal. In 1850 a process for making “coal oil” was patented in Scotland. This process was developed to augment the dwindling supplies of whale oil. The ability to produce kerosene from oil was demonstrated at Dartmouth college in 1854 and became significant with the increase in oil production in 1859, when Edwin L. Drake started drilling for oil. By 1862 kerosene from oil had completely displaced kerosene from coal.
pot-still\textsuperscript{c} to boil the crude oil so that the middle-boiling fraction (kerosene) could be collected and sold as lamp oil. Soon afterwards the internal combustion engine was invented,\textsuperscript{d} which burned gasoline instead of methane (coal gas). This created a market for the light-boiling fraction of the distillation-only refinery too. These two products, gasoline and lamp oil (kerosene), were in high demand from 1890 onwards, especially with the boom in the automobile industry after the introduction of the Ford model T in 1908.\textsuperscript{e,\textsuperscript{(2)}} As a matter of fact, world consumption of oil doubled every decade since 1880 until the Oil Crisis in 1973 (Figure 1).\textsuperscript{(3)}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Global oil consumption for the period 1860-1980.}
\end{figure}

The problem faced by refiners at the beginning of the twentieth century was that the products obtained from the distillation of crude oil were not in the same ratio as required by the market. Furthermore, as the automobile manufacturing industry got going and later the aviation industry too, improvements in engine design required a better quality fuel to deal

\textsuperscript{c} Until 1910 all crude fractionation had been done by batch distillation. Only then did continuous-operating boilers appear. The first true distillation columns were only built in the 1920's.

\textsuperscript{d} Historically it was Germany that played the key part in the development of the automobile, with the invention of the internal combustion engine by August Nikolaus Otto in 1872 and the pioneering work of Gottlieb Daimler and Karl Benz. Both Daimler and Benz realised that gasoline (petrol) would be a better fuel than gas and adapted the Otto-engine for such use. The spark-ignition system and water cooling for automobiles can both be credited to Benz (1885).

\textsuperscript{e} Most European automobiles were built for sportsmen and enthusiasts, not for the ordinary man who just needed transport. Henry Ford had the insight to provide transportation for the common man and 15 million Ford model T’s were sold between 1908 and 1927.
with the performance requirements of higher compression ratio engines. These two aspects, namely market requirements and fuel quality, were destined to become a recurring theme in crude oil refining. Upgrading crude oil by distillation only (a first generation refinery) was no longer enough and it heralded the start of the second phase in refinery development.

Markets had to be found for the heavier products from crude oil and methods had to be devised to change the properties of the products to meet the market demands. The second stage of refinery development is characterised by the introduction of reactive processes to change the nature of the crude oil. Although the development of refineries, catalysts and the chemical engineering discipline started off independently, they quickly came together to create second generation refineries. This period saw the development of many of the key technologies that are still being used in refineries today (Table 1).

Table 1. Historical development of some key refining technologies.

<table>
<thead>
<tr>
<th>Year</th>
<th>Process</th>
<th>Catalyst</th>
</tr>
</thead>
<tbody>
<tr>
<td>1920's</td>
<td>Hydrogenation</td>
<td>Ni-kieselguhr</td>
</tr>
<tr>
<td>1933</td>
<td>Paraffin isomerisation</td>
<td>AlCl₃</td>
</tr>
<tr>
<td>1934</td>
<td>Olefin oligomerisation (CatPoly)</td>
<td>Solid phosphoric acid (SPA)</td>
</tr>
<tr>
<td>1936</td>
<td>Catalytic cracking (Houdry)</td>
<td>Montmorillonite (acidic clay)</td>
</tr>
<tr>
<td>1940</td>
<td>Catalytic cracking improved</td>
<td>Amorphous silica-alumina (ASA)</td>
</tr>
<tr>
<td>1942</td>
<td>Paraffin alkylation</td>
<td>HF and H₂SO₄</td>
</tr>
<tr>
<td>1942</td>
<td>Fluid catalytic cracking</td>
<td>Amorphous silica-alumina (ASA)</td>
</tr>
<tr>
<td>1949</td>
<td>Reforming (Platforming)</td>
<td>Pt-Al₂O₃</td>
</tr>
<tr>
<td>1959</td>
<td>Hydrocracking</td>
<td>Ni/Mo- Co/Mo- silica-alumina</td>
</tr>
<tr>
<td>1964</td>
<td>Zeolite fluid catalytic cracking</td>
<td>Faujasite (Y)</td>
</tr>
<tr>
<td>1967</td>
<td>Multimetallic reforming</td>
<td>Pt/Re- Pt/Ir-Al₂O₃</td>
</tr>
<tr>
<td>1973</td>
<td>Etherification</td>
<td>Acidic resin (e.g. Amberlyst 15)</td>
</tr>
<tr>
<td>1970's</td>
<td>Zeolite based processes</td>
<td>ZSM-5, etc.</td>
</tr>
</tbody>
</table>

When the application of compression ignition engines became more widespread, markets were found for diesel fuel and fuel oil. Fuel oil started replacing coal as energy source on ships in the 1900’s.

Since the development of the compression ignition engine by Rudolf Diesel in 1893, it took some time before its use became widespread. Despite its better thermodynamic efficiency than a spark-ignition engine,
The demand for high-octane aviation gasoline during the Second World War and transportation fuel in general, also prompted technological advances in crude oil refining. Yet, crude oil was cheap and despite increasing consumption, prices were low. Many refineries did not make use of all the technologies that were on offer and the new technologies that were being developed. Second generation refineries generally used a simple topping-reforming refinery flowscheme (Figure 2), which sometimes included visbreaking too. The refineries that exploited the market for high octane gasoline during the Second World War, were left long in octane once the demand for aviation gasoline dropped in the 1950’s.

![Diagram of topping-reforming flowscheme used by most 2nd generation refineries.](image)

In the early 1970’s crude oil prices were at their lowest, with a barrel of Arabian Light crude costing only US$ 1.80. This was to change quite dramatically with the Oil Crisis at the end of 1973, when oil prices increased six-fold in a short period of time. In January 1974

compression ignition engine is heavier and more expensive, which initially made its use for automobiles less attractive, but made it very attractive for heavy vehicles and ships.

Visbreaking is a refinery process and the term is used to denote thermal cracking of residues. It is not the same as thermal cracking, which is a process used for the production of short chain olefins. The principle of operation is the same though.

A small number of very large international oil companies informally working together, controlled most of the world’s markets for oil in the 1950’s. By 1959 these companies were under economic pressure from each other and outsiders. They therefore decided to reduce their tax commitments to the producing countries to boost their profits, by posting lower prices for the crude oil on which tax was calculated. The reaction was unexpectedly strong and one of the results was the formation of the Organisation of Petroleum Exporting Countries (OPEC) at the instigation of Venezuela. During the 1960’s internal disagreements amongst member countries failed to result in any coherent collective action, but OPEC survived mostly as a result of the political solidarity felt by the Arab countries. It was this solidarity that caused the production cuts in 1973, since it was seen as a way to get back at the pro-Israel Western world after the Six Day War in 1967 and finally the Yom Kippur War that started on 6 October 1973.
a barrel of Arabian Light crude was around US$ 12, with another significant increase that were to take place at the end of the decade. Since the residue fraction of most crude oils constitute close to 50% of the total volume of crude oil, the increase in oil price not only affected the price of white products (motor-gasoline, jet fuel and diesel), but also the price of fuel oil. This resulted in a drop in the use of fuel oil and a slowdown in the growth of white product consumption. Refiners now had to produce more white products and less fuel oil, while light crude oils were becoming scarcer and more costly. This gave rise to third generation refineries that included significant residue upgrading capacity (Figure 3). Such refineries typically used topping-reforming-cracking or topping-reforming-cracking-visbreaking configurations and included a vacuum distillation column for residue fractionation.

![Diagram of Topping-reforming-cracking flowscheme with vacuum distillation and optionally visbreaking of 3rd generation refineries.](image_url)

*Figure 3. Topping-reforming-cracking flowscheme with vacuum distillation and optionally visbreaking of 3rd generation refineries.*
In the same period, the introduction of unleaded motor-gasoline was lobbied. This would remove the 5-7 octane number boost that refiners could rely on by adding tetra-ethyl lead (TEL) to the motor-gasoline. Just as the Oil Crisis forced refiners to consider crude as an expensive feedstock from which the production of high value white products had to be maximised, the need to forgo the use of TEL forced refiners to have a serious look at the quality of the products being produced. This was not an immediate concern, since lead phase-out took some time. Yet, time moved on and in the 1980’s it was realised that taking out lead was not enough, sulphur content of fuel also had to be reduced. Deeper hydrogenation of all cuts not only consumed more hydrogen, but also lowered the octane number of the motor-gasoline. This set the stage for the development of the next generation of refineries.

Refinery upgrades and new refineries planned to meet the need for high octane unleaded, low sulphur motor-gasoline had to modify their naphtha upgrading sections. It was quickly realised that paraffin quality was the key to meeting the octane demands and that isomerising C4-C6 paraffins provided a cheap and easy octane boost. It was also realised that the olefins produced by the high temperature residue upgrading units opened synthetic routes for the production of high octane motor-gasoline. Fourth generation refineries (Figure 4), typical of the 1990’s, used topping-reforming-cracking-visbreaking-alkylation-isomerisation schemes. These schemes frequently include etherification too. The number and severity of the hydrotreating operations in a fourth generation refinery are also increased, especially with respect to the refining of heavier material that is rich in sulphur.

Not all refineries have opted to change with the times and intermediate product exchange agreements are often used to balance product properties with market demands. For example, of the 99 refineries in Europe there are still 16 refineries that have not added residue upgrading capacity and are essentially still 2nd generation refineries.

---

1 The president of General Motors (GM), Mr. Cole, lobbied the introduction of unleaded gasoline. It started with a public announcement at the Society of Automotive Engineers (SAE) on 14 January 1970, which indirectly led to the Clean Air Act Amendments of 1970, signed by Pres. Nixon in November 1970. The distribution network and vehicles requiring unleaded gasoline were ready by end 1974.

2 The phase-out of lead was a gradual process and in a country like the USA that prefers motor-gasoline to diesel powered vehicles, it took 20 years to complete. By the end of the 1980’s leaded gasoline was essentially gone and by 1996 its use on highways became illegal in the USA.

3 Ironically the reduction in sulphur was called for in parallel with the phasing out of lead, but to make the introduction of unleaded gasoline a success, its impact was downplayed.

4 The three main synthetic motor-gasoline technologies based on olefins are alkylation, etherification and oligomerisation. Of these the alkylation of iso-butane with olefins is most often used. Etherification went through a boom-and-bust cycle in the USA for political reasons, but is still widely used in Europe. The use of oligomerisation is less widespread, mainly because it uses two moles of olefin per one mole of product, unlike alkylation and etherification that requires only one mole of olefin per one mole of product.
Figure 4. Topping-reforming-cracking-visbreaking-alkylation-isomerisation flowscheme, 4th generation refineries.

It will be appreciated that refineries are constantly changing and that they are subject to the same pressures that shape transportation fuel specifications. Oil refining will therefore be discussed in general terms, referring to the main separation and conversion processes needed to produce transportation fuels. This will serve as an introduction to crude oil refinery design and what factors need to be considered for grassroots refineries, including some future trends in refinery design.

2. Separation processes

Separation processes are associated with all refining units, but in the present context it refers to the primary crude oil separation that takes place at the front-end of a refinery (Figure 5).\(^{12}\) The first step is desalting, which is mainly a corrosion and fouling prevention measure. During desalting the crude oil is washed with water and caustic (NaOH) to remove dissolved salts and to convert acid chlorides, such as MgCl\(_2\) and CaCl\(_2\), to NaCl (Equation 1) that is a neutral chloride. This prevents the formation of hydrochloric acid when residual chlorides enter the refinery.
During this process some naphthenic acids are also converted to their respective carboxylate salts (Equation 2) and are removed as part of the aqueous effluent.

\[
\text{RCOOH} + \text{NaOH (aq)} \rightarrow \text{RCOONa (aq)} + \text{H}_2\text{O} \quad \cdots \ (2)
\]

This can sometimes be problematic though, because the carboxylate salts are surface active and can form stable emulsions. This hazard is reduced by coalescing and decanting the water droplets under the influence of an electric field.\(^{m,16}\)

The main separation step in any crude oil refinery is atmospheric distillation unit (ADU).\(^n\) This column fractionates the crude into straight run naphtha, kerosene, distillate and atmospheric residue. This is a crucial refining step, since it routes the molecules to the

---

\(^{m}\) A field strength in the order of 700-1000 V·cm\(^{-1}\) is typically used.

\(^{n}\) This is also referred to as the crude distillation unit (CDU).
appropriate conversion units in the refinery. The cut point of the atmospheric residue depends on the prevailing fuel specifications and crude slate used. It is typically set to satisfy the T90 specification of diesel fuel from the refinery and is usually in the range 360-375°C. The atmospheric residue is further processed in the vacuum distillation unit.

The cut points of the product fractions from the vacuum distillation unit (VDU) depend on the refinery configuration. The gas oil or vacuum distillate cuts range from 360-550°C and are typically used for lubricating oil production, but may also be converted to lighter cuts in the refinery. The heavy product from the VDU that has a boiling point above 550°C is called the vacuum residue. The vacuum residue contains most of the crude oil contaminants like metals, salts, heteroatom rich compounds and high molecular weight polycrystalline aromatic structures. The vacuum residue may be further processed in the refinery, or serve as feed for the last separation process at the front-end of the refinery, namely deasphalting.

Deasphalting is a liquid-liquid extraction whereby the last of the molecules that can be refined to white products are extracted from the vacuum residue. Light hydrocarbons, like propane, butane and pentane are typically used as solvents. The yield of deasphalted oil increases with increasing molecular weight of the solvent, but its quality decreases, while the fluidity of the asphalt is likewise decreased with increasing molecular weight of the solvent (Table 2).\(^{(12)}\)

<table>
<thead>
<tr>
<th>Properties</th>
<th>Type of solvent used for deasphalting</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Propane</td>
</tr>
<tr>
<td>Yield (mass %)</td>
<td>32</td>
</tr>
<tr>
<td>Density (kg·m(^{-3}))</td>
<td>945</td>
</tr>
<tr>
<td>Sulphur content (mass %)</td>
<td>3.8</td>
</tr>
<tr>
<td>Nitrogen content (mass %)</td>
<td>0.1</td>
</tr>
<tr>
<td>Ni+V content (µg·g(^{-1}))</td>
<td>6</td>
</tr>
<tr>
<td>Viscosity @ 100°C (cSt)</td>
<td>60</td>
</tr>
</tbody>
</table>

Dissolved gases and volatile hydrocarbons can be recovered after desalting or atmospheric distillation. The gas is typically separated into fuel gas, liquid petroleum gas (LPG) and light naphtha. The gas recovery section is used not only for the straight run gas,
but also the gas produced during conversion processes. An important component of the gas recovery section is *acid gas treatment*, which entails the removal of hydrogen sulphide (H₂S) from the gas. The H₂S is typically dissolved in a solvent from which it can be recovered and converted to elemental sulphur in a Claus or similar type of unit.¹⁷

3. **Conversion processes**

Conversion processes are required to bridge the gap between feed properties and desired product properties. Numerous refining technologies exist, but for the most part the conversion processes can be classified under the following headings:

a) **Carbon number growth.** These are technologies that increase the average molecular mass of the carbon backbone of the molecule, such as oligomerisation. Due to the heavy nature of crude oil, there is seldom use for this class of technologies in processing straight run products, but it is used in upgrading gas from cracking units by processes like alkylation.

b) **Carbon number reduction.** These are technologies that decrease the average molecular mass of the carbon backbone of the molecule, such as cracking. Many residue upgrading technologies⁰ fall into this category, because they transform molecules with a high boiling point to lower boiling products.

c) **Functionalisation.** Technologies that change the functional group to which a molecule belongs, fall in this category, like hydodesulphurisation (HDS), reforming and etherification. One could say that hydrotreating technologies result in defunctionalisation, but the removal of heteroatom-based functional groups effectively changes the class of compounds to which a molecule belongs.

d) **Rearrangement.** This is a special class of technologies that can modify the structure of a molecule, without changing the carbon number or functional group of the molecule, like hydroisomerisation and double bond isomerisation.

Some technologies perform more than one function and can be classified into more than one of these categories. This form of process intensification¹⁸ does not invalidate the

---

⁰ Residue upgrading technologies are also sub-divided into the classes “hydrogen addition” and “carbon rejection”. A secondary aim of residue upgrading is to enrich the hydrogen content of the lower boiling products. If this can be done by adding hydrogen, like in hydrocracking, which makes it a “hydrogen addition” technology, but hydrogen is a scarce commodity in a crude oil refinery. More often some of the carbon mass of the residue is sacrificed as coke or some other form of carbonaceous product to comparatively enrich the lighter products in hydrogen. Examples of such technologies are fluid catalytic cracking and coking, which make them “carbon rejection” technologies.
classification and in such cases the technology is classified according to its main aim. For example, hydrocracking is mainly a carbon number reduction technology, although it also results in heteroatom removal. This classification is important, because it describes the function of the conversion. It comes in handy to describe the type of conversion processes needed to deal with the various crude oil fractions, without immediately delving into the details associated with technology selection linked to the specifics of each refinery design.

3.1. Residue upgrading

It is appropriate to start with residue upgrading, since it is a source of lighter boiling fractions that need to be upgraded with the other straight run products. The upgrading can take two basic forms, either it aims to reduce the average carbon number of the molecules to be upgraded as transportation fuels, or it modifies the properties of the products to be used in non-fuel applications.

3.1.1. Conversion of residue to fuels

Residue upgrading technologies that have carbon number reduction as main objective are normally high temperature (>350°C) processes:

a) Fluid catalytic cracking (FCC). Typical products from FCC are gas (15-20%), naphtha in the 20-220°C boiling range (40-50%), light cycle oil in the 220-350°C boiling range (15-20%), heavy cycle oil and slurry in the >350°C boiling range (15-20%) and coke (5%).(8)

b) Visbreaking. Typical products from visbreaking are gas (1-5%), naphtha in the 20-165°C boiling range (5-10%), gas oil in the 165-350°C boiling range (10-25%) and residue in the >350°C boiling range (60-85%).(19)

c) Coking. Typical products from delayed coking are gas (5-10%), naphtha in the 20-195°C boiling range (10-25%), gas oil in the >195°C boiling range (30-65%) and coke (10-35%).(20)

d) Hydrocracking. Typical products from hydrocracking are gas (5%), naphtha in the 20-180°C boiling range (15-20%), kerosene in the 180-250°C boiling range (30-40%), gas oil in the 250-360°C boiling range (25-45%) and residue in the >360°C boiling range (0-20%).(7)
With the exception of hydrocracking, all the other technologies are carbon rejection processes. They also produce olefins, which is one of the main hydrocarbon classes required in naphtha upgrading, but that is not present in crude oil.

3.1.2. Non-fuels application of residues

Residues are not necessarily upgraded by conversion technologies. Some of the main non-fuel related applications of residues rely mostly on separation processes to achieve the desired product characteristics.\textsuperscript{(21)}

a) **Lubricants.** There are numerous special lubricant types and applications, but most lubricants can be classified as either oil or grease. Lubricating oil is generally aliphatic in nature, with solvent extraction of the aromatics being used to improve the viscosity index and solvent dewaxing being used to remove linear paraffins to improve the cold flow properties. Lubricating grease is a semi-solid material that is obtained by dispersing a gelling agent in a lubricating oil.

b) **White oils.** This is a term used for highly refined lubricating oils, principally where the oil has been significantly dearomatised. These white oils were previously made from light oil cuts treated with oleum to remove the aromatics, but presently deep catalytic hydrodearomatisation (HDA) is more often used.

c) **Paraffins and waxes.** These are by-products from solvent dewaxing and are used in numerous applications ranging from food packaging to polishes and cosmetics.

d) **Bitumen.** There are many grades of bitumen that differ in hardness and flow properties. It can be produced as vacuum residue or by deasphalting of vacuum residue. Air oxidation, typically at temperatures higher than 200°C, is often employed to modify the structure and to produce hard grades of bitumen. By increasing the oxidation temperature, air contact time or air pressure the hardness can be increased.\textsuperscript{(22)} Applications of bitumen include waterproofing, fillers, insulation materials and road binders. Cracked residue does not make a good feed material for bitumen production.\textsuperscript{(23)}

e) **Coke.** Although coking has been discussed as a conversion process to produce fuels, it also produces coke. The quality of the coke determines its application possibilities, for example graphite, pigments, electrodes or reducing agents in the metallurgical industry.

3.2. Diesel and jet fuel upgrading
Diesel and jet fuel have many specification characteristics in common and will therefore be treated together. The straight run kerosene and distillate have the right carbon number distribution and mainly requires a reduction in sulphur content and aromatic content (functionalisation) to meet fuel specifications. This can be achieved by hydrotreating.\textsuperscript{p}

Depending on the crude oil, the product might still contain linear paraffins, which is good for diesel cetane, but could be detrimental to diesel cold flow properties and cause the jet fuel to fail the freezing point specification. The linear paraffins can either be modified by hydroisomerisation (rearrangement), or the linear paraffins can be removed by solvent dewaxing. The other fuel properties that may be lacking can be fixed by fuel additives.

It is clear that the upgrading of crude oil derived straight run kerosene and distillate is not complicated. Since the requirements for jet fuel and diesel are almost the opposite of that required for motor-gasoline, it is anticipated that naphtha upgrading will be much more involved.\textsuperscript{q}

### 3.3. Naphtha and gas upgrading

The upgrading of naphtha and gas to motor-gasoline is unit-wise by far the most intensive refining operation. The main adjustments that have to be made to the properties of the fuel are related to the sulphur content and octane number (functionalisation and rearrangement), as well as the average molecular mass when dealing with gas (carbon number growth). Getting rid of the sulphur is not difficult, since naphtha range material contains mostly mercaptans and other easy to remove sulphur containing compounds, which can be removed by mild hydrotreating or sweetening. Meeting the octane number specification is more difficult. Crude oil derived straight run naphtha typically has a low octane number (LSR from Arabian Light crude has a RON of about 60 and the heavy naphtha has a RON of less than 25)\textsuperscript{16} and increasing the octane number requires a complex refinery scheme.

The main sources of octane are aromatics and oxygenates (ethers and alcohols), but the volume of these compounds are limited by motor-gasoline specifications. Roughly half

\textsuperscript{p} Hydrotreating to meet diesel fuel specifications will automatically ensure that the mercaptan specification of jet fuel is met. Previously this would not necessarily have been the case and the kerosene cut would have to go through a sweetening process.

\textsuperscript{q} It may seem that the increased refining complexity required for motor-gasoline production compared to diesel fuel production is another reason to nudge the market in the direction of increased diesel fuel use. Surprisingly, from a refining perspective this is not true. Catalytic reforming is the main source of refinery hydrogen and is linked to motor-gasoline production. Furthermore, since transportation fuels are sold by volume, motor-gasoline effectively commands a higher price than diesel fuel on account of its lower density.
of the motor-gasoline has to be paraffinic and it is the quality of the paraffins in the fuel pool that determines the refining needs (as discussed in Chapter II). As a rule of thumb, the required minimum octane number of the paraffins can be calculated by a simple formula (Equation 3) based on the octane specification ($RON_{spec}$) and the volume fractions ($V_i$) and octane numbers ($RON_i$) of the various compound classes:

$$RON_{paraffin} \geq \frac{[RON_{spec} - (RON\cdot V)_{aromatic} - (RON\cdot V)_{oxygenate} - (RON\cdot V)_{olefin}]}{V_{paraffin}} \quad \ldots (3)$$

Most modern crude oil refineries use catalytic reforming.\textsuperscript{5} The reformer is not only the main source of aromatics for the motor-gasoline, but also the main source of hydrogen in the refinery. The octane number of aromatic rich reformate can be manipulated by varying the severity of the operating conditions of the reforming unit, but in practise reformate generally has a RON of 96-102, with an aromatic content of around 70%.\textsuperscript{9}

Not all refineries have etherification units and with present pressure on fuel ethers in the USA, many refineries are forced to use ethanol instead. Nevertheless, ethanol has a RON of 120,\textsuperscript{24} while the typical fuel ethers have a slightly lower octane number. Olefin oligomerisation, which typically yields a RON of about 96, is practised in some refineries, but FCC derived olefins are more often used for etherification and aliphatic alkylation.

Using Euro-4 specifications as reference, aromatics can be blended to 35%, olefins to 18% and ethers and alcohols to 10-15%. If typical, realistic refinery values are substituted into Equation 3,\textsuperscript{6} the paraffin RON required to make 95 octane motor-gasoline can be calculated for scenarios with and without oxygenates and olefins (Equations 4-7):

$$RON_{paraffin} \geq \frac{[95 - (98\cdot 0.5)_{aromatic} - (120\cdot 0.1)_{oxygenate} - (96\cdot 0.15)_{olefin}]}{(0.25)_{paraffin}} \geq 79 \quad \ldots (4)$$

$$RON_{paraffin} \geq \frac{[95 - (98\cdot 0.5)_{aromatic} - (120\cdot 0.1)_{oxygenate} - (0)_{olefin}]}{(0.4)_{paraffin}} \geq 85 \quad \ldots (5)$$

$$RON_{paraffin} \geq \frac{[95 - (98\cdot 0.5)_{aromatic} - (0)_{oxygenate} - (96\cdot 0.15)_{olefin}]}{(0.35)_{paraffin}} \geq 91 \quad \ldots (6)$$

$$RON_{paraffin} \geq \frac{[95 - (98\cdot 0.5)_{aromatic} - (0)_{oxygenate} - (0)_{olefin}]}{(0.5)_{paraffin}} \geq 92 \quad \ldots (7)$$

It is clear that even when oxygenates and olefins are added to the fuel, some paraffin upgrading is required to increase the octane number of the straight run naphtha. The

\textsuperscript{5} A catalytic reformer will typically not be found in refineries that do not produce on-specification motor-gasoline as a final product. In such refineries hydrogen is produced by other means, for example steam reforming of methane.

\textsuperscript{6} Since reformate is not a pure aromatic stream, but contains about 70% aromatics at a RON of 98, 50% of the motor-gasoline can be made up of reformate without violating the 35% maximum aromatic specification.
following upgrading pathways can be considered, assuming that a source of olefins is available from residue conversion units in the refinery:

a) *Etherification.* Depending on the branched olefins being available and the preferred alcohol, the typical fuel ethers produced are 2-methoxy-2-methylpropane (MTBE, methyl tertiary butyl ether), 2-ethoxy-2-methylpropane (ETBE, ethyl tertiary butyl ether) and 2-methoxy-2-methylbutane (TAME, tertiary amyl methyl ether). These ethers all have a RON in the range 115-118.(24) It is also possible to directly etherify light FCC gasoline,(25)(26) yielding a product with a calculated blending RON of 110.(27) However, it has been shown that most C₆ and heavier ethers have poor octane numbers, unless the source olefin is highly branched.(28)

b) *Olefin oligomerisation.* There are various technologies available for olefin oligomerisation,(29) but when the focus is on unhydrogenated motor-gasoline, the octane values from many processes fall in a narrow band around RON 92-97 and MON 79-82. Olefinic motor-gasoline is therefore close to octane neutral, albeit somewhat MON constrained.

c) *Aliphatic alkylation.* The alkylation of iso-butane with olefins to produce alkylate, is often used as a source of high octane paraffins; RON 92-97 and MON 90-94.(30)(31)(32) Although the product is highly desirable, current commercial processes are all based on either sulphuric acid (H₂SO₄), or hydrofluoric acid (HF). Much effort has been spent on the development of more environmentally friendly solid acid catalysed processes, but with limited success.(33) It can be argued that aliphatic alkylation is essential to meet the requirements for high octane paraffins, but environmental concerns keep looming in the background.

d) *Hydroisomerisation.* The skeletal isomerisation of paraffins in the C₄-C₆ range is often practised. Butane isomerisation typically provides a source of iso-butane for aliphatic alkylation when it is used in the refinery, while hydroisomerisation of LSR (C₅-C₆) is routinely employed to produce an isomerate with RON 78-92, depending of the recycle configuration of the technology.(10) The hydroisomerisation of C₇ and heavier fractions is actively researched, but has met with little success.(34)(35)

It can therefore be said that all naphtha and gas upgrading sections of a crude oil refinery has a hydrotreater to remove sulphur and reformer to produce aromatics. In addition to these units, one or more paraffin upgrading units needs to be present, namely aliphatic alkylation and hydroisomerisation, with optionally etherification and olefin oligomerisation.
An external source of alcohols is required to produce motor-gasoline blends that require oxygenates.

4. Future crude oil refineries

4.1. Change drivers in crude oil refining

Based on the historical development of crude oil refineries, it does not take a huge leap of faith to predict that crude oil refineries will continue to change. The drivers for change are inherent to the purpose of a refinery, namely to transform a raw material into a more valuable end product that meets market demands. These are:

a) Crude oil availability and cost. Although it is unlikely that crude oil will run out in the short term, it is a finite resource and some of the more sought after types of crude oil may become scarcer more quickly. This underpins two threats to a refinery, namely the availability and the cost of the raw material.

b) Refinery capital and operating cost. To turn the raw material into a more valuable product, the market must be willing to buy the product at a higher price than the cost of upgrading it. To process cheaper more readily available raw materials (lower quality heavy crudes), will require capital to be invested in residue upgrading units and result in an increase in the operating cost. Refiners are also under pressure to keep their environmental footprint small and processing heavier crudes is environmentally speaking a move in the wrong direction. The decision to modify a refinery to deal with cheaper difficult crudes, is therefore not such a clear-cut investment decision. Furthermore, there is a rising cost associated with environmental conservation and the refinery upgrades needed to keep within legislated emission levels. The underlying threat is that more efficient, more environmentally friendly or even alternative technologies for the production of transportation fuel may come along to undermine the profitability of investments in refining infrastructure. Conversely there is the threat that in times of high profit, governments might impose a “windfall” tax on refiners, as has indeed happened in the past.

c) Market demands. To sell transportation fuel, the fuel must comply with the fuel specifications that have been changing to require increasingly refined products. This places strain on the refiner, because the environmental concerns giving rise to more stringent fuel
specifications, translate into a larger environmental footprint of the refinery. This reality is not always appreciated by the politicians and the lawmakers. Furthermore, the market for refined products is not homogeneous and grows at different rates in different regions. In general market demand for fuel oil is decreasing and market demand for white products are increasing, but this says nothing about the substructure of white product demand. Product price also affects the overall market demand and may selectively suppress the use of some fuel types when the price is too high, or stimulate their demand when the price is low. This gives rise to fears about over-investment in refining capacity, as well as under-investment in refinery flexibility to deal with market swings.

4.2. Changes in crude oil refining

The impact of the change drivers will be seen both in refinery design, as well as in the transportation fuel industry. In refineries, effects will most likely be seen in the following areas:\(^{40}\)

a) Residue upgrading. More efficient residue upgrading is key to refining economics. This is driven by the lower demand for heavy products like fuel oil and the price differential between expensive light crude oils and cheap heavy crude oils.\(^{(3)(12)(41)}\) The need for molecules from residue upgrading units, especially olefins,\(^{(42)}\) is also becoming more apparent with the need for high octane paraffinic blending components for motor-gasoline production. This further emphasises the importance of residue upgrading in future. The nature of the residue upgrading capacity that is added has one natural side-effect, namely that it influences the motor-gasoline to distillate ratio produced by the refinery. In Europe this is a serious concern and may cause the significant swing from motor-gasoline to diesel to slow down and reach a natural refinery driven equilibrium.\(^{u(15)}\) Any technology breakthrough in residue upgrading technology, for example novel technology to improve or supersede FCC, will have a tremendous impact on future crude oil refinery flowschemes.

b) Hydrogen production. The drive to produce more ultra low sulphur transportation fuels will increase the need for hydrogen in the refinery. The increase in hydrogen demand is exacerbated by the lowering of the maximum aromatics content in fuel, which decreases the

---

\(^1\) In many European countries the cost of compliance to environmental legislation increased between 400-2500% from 1985 to 1993.

\(^u\) In Europe many refiners are already having difficulty meeting diesel fuel demand and quality. This problem is most acute in refineries having FCC’s (57 refineries), which represent 76% of the European refining capacity. It is not clear to what extent the political landscape will change to cope with this challenge.
hydrogen production in the refinery. Additional hydrogen will also be required if residue upgrading increasingly employs hydrogen addition, rather than carbon rejection. This may necessitate investment in on-purpose hydrogen production units. At this stage hydrogen production is costly and chances of a “hydrogen economy” bleak, but advances in technology may make refineries more hydrogen rich in future.

c) Refinery complexity and cost. As fuel specifications drive the fuel composition further away from the crude oil composition, more units will be required to make final products. This has implications for refinery complexity and cost, increasing the minimum refinery size that makes economic sense. To offset the cost, refineries may increasingly opt for co-generation of power, as well as to co-produce chemicals. With increasing focus on residue upgrading and hydrogen production, chemicals like propylene from FCC and aromatics from reforming are natural choices for co-production.

d) Renewable fuel. If the inclusion of bio-derived components in transportation fuel becomes mandatory, it will be impossible for a crude oil refinery to produce on-specification fuel with only crude oil as raw material. At present, most fuels that include a bio-derived component are only blends, but there are already investigations dealing with possible co-processing.

e) Synthetic fuel. The need for hydrogen in a refinery can be addressed by investing in on-purpose hydrogen production, which may entail gas reforming (water-gas shift). This would not only make hydrogen available, but could potentially also make synthesis gas (H₂ + CO) available to the refinery. If this happens, crude oil refineries may in future become integrated with synthesis gas based synthetic fuel technologies, such as Fischer-Tropsch and Syngas-to-methanol. The benefit of CO₂ capture to enhance oil production in addition to the use of Fischer-Tropsch technology to produce fuels, have also been advocated. Although it should be noted that the possibility of using CO₂ for enhanced oil recovery, is very location specific. Furthermore, gasification technology could be a convenient sink for excess coke production from residue upgrading by delayed coking, since the coke can be employed as gasifier feedstock. This approach has previously been explored in the Exxon Donor Solvent (EDS) process for direct coal liquefaction. The EDS process

---

v It is interesting to speculate on the political ramifications of this situation. Big oil companies may want to buy into agriculture, which is arguably then an essential part of their business. Since bio-fuels are subsidised, the oil companies would be subsidised, an issue that is already boiling up with the 2007 Tyson-ConocoPhillips deal. According to this deal, a “renewable” diesel fuel will be produced from animal fat by co-processing the animal fat with crude oil derived hydrocarbons in a thermal depolymerisation (cracking) unit.
was combined with the gasifier section from an Exxon Flexicoker\(^{(20)}\) to convert the coal residue into hydrogen, which is needed for coal liquefaction.

### 4.3. Future crude oil refineries

Translating trends in crude oil refining into a futuristic flowscheme is fraught with the same obstacles encountered in devising any generic refinery flowscheme. Views on future crude oil refineries should therefore not be seen as a concrete entity, but rather as an expression of philosophy, with very different pictures emerging.

A fairly traditional approach is presented by Heinrich from the Institut Français du Pétrole (Figure 6).\(^{(12)}\) The intensity of residue upgrading and the need for additional hydrogen in the refinery is reflected in the addition of a hydrocracker and partial oxidation unit. Although it is not explicitly indicated, the partial oxidation unit is coupled with some form of syngas-to-methanol technology, making the refinery self-sufficient in terms of methanol for etherification. A lot of emphasis has been placed on etherification for the production of high octane oxygenates, a view that is questionable in USA context, but probably still valid for Europe. The appearance of an olefin oligomerisation unit is
interesting, since it implies a much more olefin rich refinery. The increase of olefins is a consequence of the intensification of residue upgrading, but also a means to reduce the hydrogen requirements. Of the five trends identified in the previous section, only the inclusion of renewable fuel is not reflected and it can be argued that it shouldn’t be, since it does not require co-refining, only blending.

A much more radical approach, focussing on the environmental impact of the refining units, is presented by Chen from ExxonMobil (Figure 7). The flowscheme is not always technology specific, with technology classes being grouped together, like olefin upgrading and deep hydroprocessing. This hides the increased complexity of the refinery. The intensity of residue upgrading and hydrogen production is reflected by the use of a thermal cracker, fluid coker, high temperature FCC and gasifier. The gasifier also enables the production of synthetic fuels by other means, like Fischer-Tropsch. In general there is a clear fuel synthesis focus and “the heart of this new scheme resides in olefin upgrading technologies”. The

---

If C₈ is taken as a representative carbon number for motor-gasoline, the ability to include octenes (C₈H₁₆) from olefin oligomerisation, rather than octanes (C₈H₁₈) in the fuel pool, saves about 10% hydrogen for that fraction.
exclusion of aliphatic alkylation and hydrocracking is interesting. The exclusion of the former is understandable from an environmental perspective and the latter is excluded purely as result of the proposed configuration. Again four of the five refining trends are embodied in the flowscheme, with only renewable fuel not being shown.

The consequence of the trends in crude oil refining is that future crude oil refineries will become more hydrogen and olefin rich and have the capacity to integrate synthesis gas based synthetic fuel. It is also expected that peak oil production, that is likely to be reached soon,\(^{(53)(54)}\) will have a significant impact on transportation fuel production. The emergence of hybrid crude oil refineries that co-process alternative materials like tar sands or coal might fundamentally change the way in which crude oil refining is approached in future.

5. Literature cited

(13) Odell, P. R. Oil and world power. Background to the Oil Crisis, 3rd ed.; Taplinger: New York, 1975.


(36) Snow, N. Are upgrades to process lower-quality crudes worth the investment? World Refining 2004, 14:8, 52.


(41) Higgins, T. Residual fuel will continue to drive refining needs. World Refining 2004, 41:2, 4.


