Tikrit University

The College of Petroleum Processes Engineering

Petroleum Systems Control Engineering

Department

Petroleum Refining Processes

Fourth Class

Lecture 13

By

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Alkylation Process

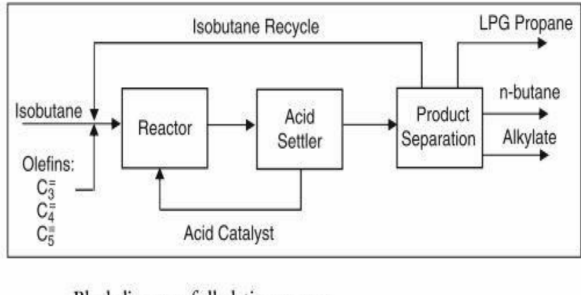
Alkylation is the process of producing gasoline range material light olefins (primarily propylene and butylene) with isobutane in the presence of a highly acidic catalyst, either

sulfuric acid or hydrofluoric acid. The product (alkylate) contains a mixture of highoctane, branched-chain paraffinic hydrocarbons. Refinery gases produced from different units are collected and sent to the gas plant. Olefins and isobutanes are separated and used as a feed to the alkylation plant to produce gasoline which can be sent to the gasoline pool.

Alkylation Processes

Alkylation is catalyzed by a strong acid, either sulphuric (H2SO4) or hydrofluoric (HF). In the absence of catalysts, alkylation between isobutene and olefin must be run under severe conditions such as T = 500 °C (932 F) and P = 200-400 bars (2940–7080 psia). In the presence of an acid catalyst, the reaction temperature will be lower than 50 °C (122 F), and the pressure will be lower than 30 bars (441 psia). The major difference in using either acid is that isobutane is quite insoluble in H2SO4 but reasonably soluble in HF. This requires the use of high isobutane/olefin ratios to compensate for low solubility in H2SO4. Furthermore, the reaction must occur at low temperature. The alkylation process consists of running the hydrocarbons in liquid form (enough pressure is used to ensure that) and at low temperature and with a high isobutane (iC4) to olefin (such as C4) ratio. The reaction products are sent to an acid settler where the acid is recycled back to the reactor. Products are then separated into

gaseous LPG propane and n-butane and the desired product of alkylate. A block diagram of the process is shown in Figure below:

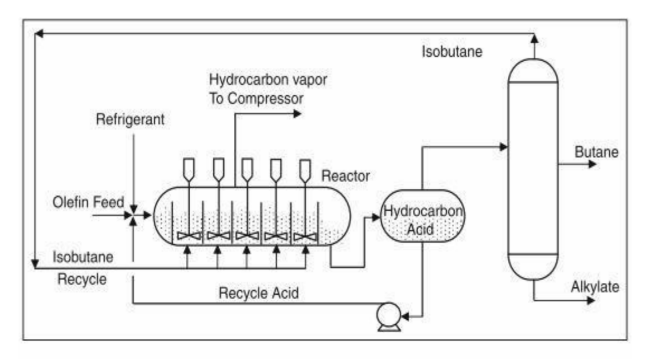


Block diagram of alkylation process

Process Flow: Sulfuric Acid Alkylation

In sulfuric acid (H_2SO_4) alkylation units, the feeds – propylene, butylene, amylene, and fresh isobutane – enter the reactor and contact sulfuric acid with a concentration of 85 to 95%. The reactor is divided into zones. Olefins are fed through distributors to each zone, and sulfuric acid and isobutanes flow over baffles from one zone to the next.

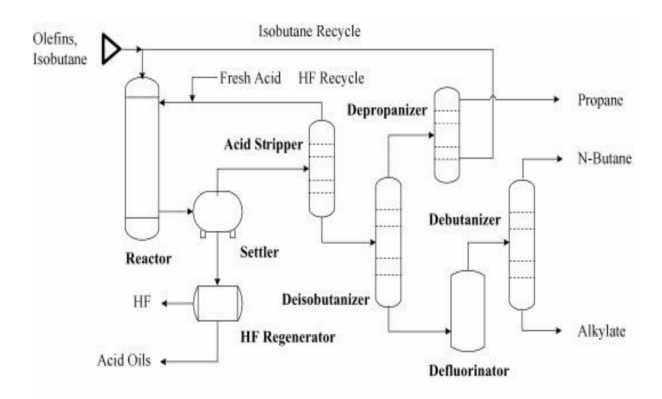
The reactor effluent goes to a settler, in which hydrocarbons separate from the acid. The acid is returned to the reactor. The hydrocarbons are washed with caustic and sent to fractionation. The fractionation section comprises a depropanizer, a deisobutanizer, and a debutanizer. Alkylate from the deisobutanizer can go directly to motor-fuel blending, or it can be reprocessed to produce aviation-grade gasoline. Isobutane is recycled. A block diagram of the process is shown in Figure below.



sulphuric acid alkylation process

Hydrofluoric Acid Alkylation

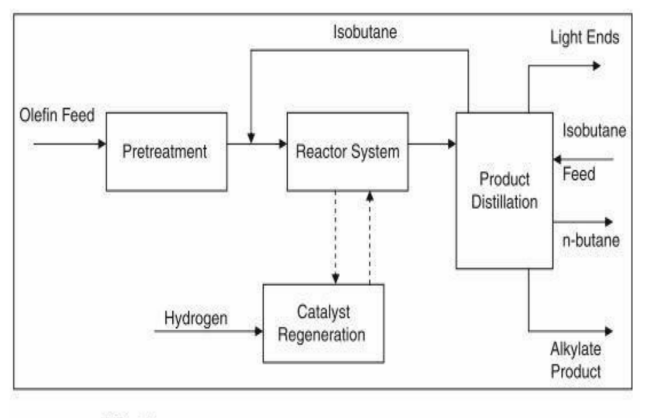
Figure below shows a process schematic for hydrofluoric acid (HF) alkylation. Olefins and isobutane are dried and fed to a reactor, where the alkylation reaction takes place over the HF catalyst. The reactor effluent flows to a settler, where the acid phase separates from the hydrocarbon phase. The acid is drawn off and recycled. The hydrocarbon phase goes to a deisobutanizer (DIB). The overhead stream, containing propane, isobutane, and residual HF, goes to a depropanizer (DeC3). The DeC3 overhead goes to an HF stripper. It is then treated with caustic and sent to storage. Isobutane from the DIB main fractionator is recycled. The bottom stream from the debutanizer goes to product blending.



HF alkylation

AlkyClean Process

Lummus technology has developed a solid acid catalyst gasoline alkylation technology. The AlkyClean process employs a zeolite catalyst coupled with a novel reactor processing to yield a high quality alkylate product. The process shown in Figure below consists of four main sections: feedstock pretreatment, reaction, catalyst regeneration and product distillation. An olefin feed is preheated and fed with the isobutane recycle to the reactor. The reactor operates at 50–90 °C (122–194 F) with liquid phase conditions. Multiple reactors are used to allow for the catalyst regeneration cycle. During regeneration of dissolved hydrogen while maintaining liquid phase alkylation reaction conditions. This minimizes energy consumption during the switching of the operation. The swing reactor coupled with long catalyst life allows the refiner to work without the need of taking the reactor off-line for moderate temperature regeneration that restores the catalyst activity completely.



AlkyClean process

Effect of Operating Conditions

Olefin Type

The presence of propylene or pentene with butane will lower the octane number and increase the acid consumption. The octane number of alkylates produced from light olefins is given in Table below:

Types of Olefin	RON		MON	
	HF	H ₂ SO ₄	HF	H ₂ SO ₄
Propylene	91-93	91-92	89-91	90-92
Butene-1	90-91	97-98	88-89	93-94
Butene-2	96-97	97-98	92-93	93-94
Isobutene	94-95	90-91	91-92	88-89
Amylene	90-92	91-92	88-89	89-91

Effect of type of olefin on alkylate octane number

Isobutane Concentration

The $iC_4/C_4^=$ ratio has an important role regarding the quality of alkylate produced and the amount of sulphuric acid consumption. The following reasons explain the behaviour.

- High isobutane concentration ([iC₄]) prevents olefin polymerization which results in low quality alkylate and high sulphuric acid consumption.
- Solubility of *i*C₄ ≪ C⁼₄. Thus high a concentration of *i*C₄ is required in the mixed hydrocarbons to compensate for its low solubility.
 - The rate of alkylate formation increases while the rate of formation of undesirable heavy alkylates decreases as *i*C₄ increases, as will be discussed later.
 - As isobutane increases, alkylate MON increases and sulphuric acid consumption decreases.

For all these reasons, the $iC_4/C_4^=$ ratio is kept in industrial operation between 5:1 and 15:1 as the external isobutane to olefin (I/O) ratio. Inside a reactor with high circulation, this ratio becomes 100–1000:1.

Acid Strength

An optimum value of acid strength of 90 wt% H2SO4 is maintained by adding fresh concentrated acid (98–99 wt%). As the strength of the acid decreases, the acid consumption increases with the octane number decreases. The minimum acid strength required to operate the system should not be lower than 85 wt%. To provide a sufficient margin of safety, acid strength is kept around 90 wt%. The acid strength decreases because of the formation of gums and other products resulting from the reaction with

other impurities. Thus, acid makeup has to be added.

Degree of Agitation

When the hydrocarbons are dispersed in sulphuric acid, the speed of the impeller determines the dispersed phase size (droplet diameter) and hence, the interfacial contact area. The reaction rate of iC4 and C4 is quite fast, and the reaction is controlled by mass transfer.

Space Velocity

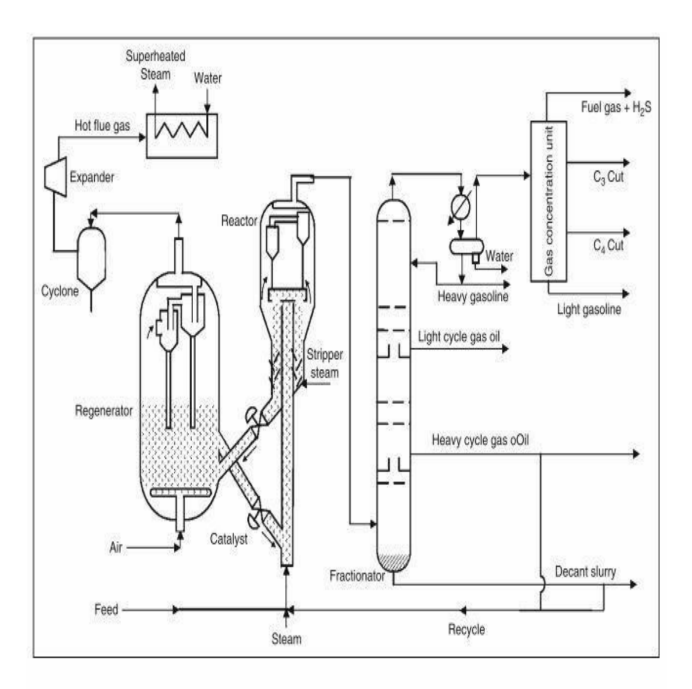
The olefin space velocity is defined as:

 $(SV)_{o} = \frac{Olefin \text{ volumetric rate (bbl/h)}}{Acid \text{ volume in contactor (bbl)}}$

The residence time in the reactor is (1/(SV)o) and is defined as the residence time of the fresh feed and externally recycled isobutane in the reaction mixture. Since the alkylation reaction is very fast, the residence time is not a limiting parameter. However, as the space velocity increases, the octane number tends to decrease while acid consumption tends to increase. Residence time for sulphuric acid is usually from 5 to 40 min, and for hydrofluoric acid, it is 5–25 min.

Reaction Temperature

The reaction thermodynamics and kinetics are favoured at low temperatures. Sulphuric acid alkylation units are operated at 5–10 °C (40–50 F). Above 10 °C, oxidation and side reactions are promoted, and the deteriorate-alkylate yield and quality while acid consumption increases. It is impossible to run the reaction below 0 °C (32 F) because acid viscosity will be too high and agitation becomes difficult. Above 21 °C (70 F), the polymerization of olefin will occur, and the octane number of alkylate decrease. For HF alkylation, the reaction temperature is less significant and is between 21 and 38°C (70 and 100 F).



Fluid catalytic cracking process flow sheet