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(54) PARAFFIN ALKYLATION PROCESS

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- (*) Notice: Subject to any disclaimer, the term of this OTHER PUBLICATIONS
patent is extended or adjusted under 35 albright Tyle F. Kranz, Ken F. Alkylation
- $1992, 91, 100$ Appl. No.: $11/402,079$ * cited by examiner
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- (52) U.S. Cl 585/716:585.7717: 585/719 ent from the first alkylation system is fed to an interim debu
- 585/717, 719

See application file for complete search history.

U.S. PATENT DOCUMENTS

3.236,912 A * 2/1966 Phillips 585,331 24 Claims, 4 Drawing Sheets

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patent is extended or adjusted under 35 Albright, Lyle F., Kranz, Ken E., Alkylation of Isobutane with U.S.C. 154(b) by 189 days. Pentenes Using Sulfuric Acid as a Catalyst..., Ind. Eng. Chem. Res. 1992, 31, 475-481, American Chemical Society.

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US 2007/0238912 A1 Oct. 11, 2007 (57) ABSTRACT

(51) Int. Cl. $C07C2/58$ (2006.01) A process for the alkylation of isobutane is disclosed wherein $C07C2/58$ isobutane is fed to two separate alkylation systems. The efflu-
ent from the first alkylation system is fed to an interim debu-(58) Field O f Classification search s s 585/716 tanizer where the C's are separated from the alkylate prod uct. The overhead C product is then fed to the second alkylation system to provide the isobutane. The effluent from the second alkylation system is fed to a traditional deisobu (56) References Cited tanizer to prevent any build up of normal butanes in the system.

FIG. 1

FIG. 2

FIG. 3

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PARAFFIN ALKYLATION PROCESS

BACKGROUND OF THE INVENTION

1. Field of the Invention

The present invention relates to the alkylation of paraffinic hydrocarbon feed stocks wherein an olefin is reacted with isobutane to produce an alkylate product. More particularly the invention relates to a process wherein an intermediate debutanizer is utilized between two alkylation systems to 10 provide a first alkylate product and a concentrated isobutane stream for use in a the second alkylation system. The process eliminates the need for a second deisobutanizer.

2. Related Information

Alkylation is the reaction of aparaffin, usually isoparaffins, 15 with an olefin in the presence of a strong acid which produces paraffins, e.g., of higher octane number than the starting materials and which boil in range of gasolines. In petroleum refining the reaction is generally the reaction of a C_3 to C_5 olefin with isobutane.

In refining alkylations, hydrofluoric or Sulfuric acid cata lysts are most widely used. For sulfuric acid catalyzed alky lation low temperature or cold acid processes are favored because side reactions are minimized. In the traditional pro carbon reactants are dispersed into a continuous acid phase. In view of the fact that the cold acid process will continue to be the process of choice, various proposals have been made to improve and enhance the reaction and, to some extent, mod erate the undesirable effects. cess the reaction is carried out in a reactor where the hydro- 25 ization-alkylation process.

SUMMARY OF THE INVENTION

Briefly the present invention is an alkylation process which utilizes an interim debutanizer or stripper between at least two 35 or more alkylation systems, preferably a process for the alky lation of isobutane with an olefin comprising a first and sec ond alkylation systems wherein the effluent from the first alkylation system is fed to a debutanizer to produce an over head and a first alkylation product, and wherein the overhead 40 is fed to the second alkylation system. The capital investment benefits and energy consumption benefits, provided by the new processing scheme stem from the ability of having or producing a predominantly n-butane free olefin feedstock to be used in a first alkylation system or unit. This allows for 45 significantly increased alkylation production without the tra ditional expense of additional deisobutanizer capacity, while still maintaining high quality motor-fuel alkylate production.
As such, it provides refiners a new option for retrofitting As such, it provides refiners a new option for retrofitting existing equipment for use in expanding their facility alkyla tion capacity.

Two general process schemes are presented. The first general process scheme involves the use of two alkylation sys tems and requires an oligomerization reactor and a smaller distillation or stripping column for separation of C_4 's from 55 heavier C_5 + materials. The first option can allow for significant reuse of equipment previously utilized for MTBE pro duction. The second major scheme allows for processing either C_3 's or C_5 's in a first alkylation system with an interim debutanizer and processes C_4 's in a secondary alkylation ω_6 Alkylation With C_4 Olefin Feedstock system with a deisobutanizer. Several variations to the two general process schemes exist, especially for cases where alkylation of C_3 through C_5 olefin containing feedstocks is desired.

The key nerem, to be able to off-load the need for addi- 65 tional deisobutanizer capacity, is in obtaining a nearly n-bu tane free olefin feed stream which may be used during an

intermediate alkylation stage. This eliminates the majority of n-butane to this intermediate alkylation stage which in turn eliminates the need for an additional isobutane/n-butane frac tionation step (deisobutanizer).

In one embodiment the process for the alkylation of isobu tane with an olefin comprises:

(a) feeding a first stream containing isobutane and a second stream containing an olefin, preferably an FCC C_4 stream containing normal butene, isobutene, normal butane and isobutane; or a mixture of: C_4 and C_5 olefins; C_3 and C_4 olefins; or C_3 , C_4 and C_5 olefins, to a first alkylation system where a portion of the isobutane is reacted with a portion of the olefin to form an alkylate containing stream;

(b) feeding the effluent from the first alkylation system to a debutanizer where C_4 's are removed as a first overheads and a first alkylate product is removed as a first bottoms:

(c) feeding the first overheads and a third stream containing isobutane to a second alkylation system where isobutane is reacted with C_4 olefins in said first overheads to form a second alkylate product.

BRIEF DESCRIPTION OF THE DRAWING

FIG. 1 is a simplified flow diagram of a basic C4 oligomer-

FIG. 2 is a simplified flow diagram for the alkylation of isobutane with C4 and C5 olefins.

FIG. 3 is a simplified flow diagram for the alkylation of isobutane with C3, C4 and C5 olefins.

FIG. 4 is a plot of alkylate quality as a function of isopen tane in the feed.

FIG. 5 is a plot of isopentane yield as a function of isopentane in the feed.

DETAILED DESCRIPTION OF THE INVENTION

As herein defined an individual alkylation system com prises all the necessary equipment for production of a dilute alkylate stream from an olefin containing stream and an isobutane rich stream. Sulfuric acid, hydrofluoric acid and solid acid catalyst alkylation processes are contemplated. Such systems are well known in the art.

Olefin feed streams utilized herein may be hydrotreated to remove dienes prior to entering the alkylation process. The removal of butadienes and pentadienes is an important ele ment in increasing catalyst life in the alkylation process. For processes which include oligomerization reactors to provide a purified oligomer product, the hydrotreatment step may be included (using reactive distillation) in the distillation col umn used to produce the oligomer stream. For feeds to oli gomerization reactors, the removal of basic compounds is required to maintain catalyst life. For butylene containing streams this may include the use of a water wash column for removal of nitriles. Catalysts used in oligomerization reactors may be acid resins, such as Amberlyst 15 or related oleum derived resins and may include phosphoric acid derived cata lysts, such as those known to the industry as SPA (solid phosphoric acid) catalysts.

In this case the means for obtaining the high purity olefin feed from a fluid catalytically cracked (FCC) C_4 feedstock is through the step of oligomerization, which is characterized as a process for the alkylation of isobutane with butenes con tained in an FCC C_4 stream comprising the steps of:

(a) feeding a first stream comprising an FCC C_4 stream containing normal butenes and isobutenes to an oligomeriza

tion reaction wherein the isobutenes react with each other to produce an effluent containing C_5 and higher oligomers and normal butenes;

(b) fractionally distilling the effluent from the oligomer ization reaction wherein the C_4 and lighter material is sepa- 5 rated as a first overheads from the C_5 and heavier material as a first bottoms;

(c) feeding the first bottoms and isobutane to a first alky lation Zone to produce a first alkylate stream containing alky late and unreacted C_4 's;

(d) feeding the first alkylate stream to a debutanizing Zone wherein the unreacted C_4 's are removed as a second overheads and alkylate product is removed as a second bottoms;

(e) feeding the first overheads and isobutane to a second alkylation Zone wherein the olefins in the first overheads are reacted with isobutane to produce a second alkylate stream containing alkylate, unreacted isobutane and unreacted nor mal butane;

(f) feeding the second alkylate stream to a deisobutanizing $_{20}$ Zone wherein isobutane is removed as a third overheads, alkylate is removed as a third bottoms and normal butane is removed as a side stream.

Optionally, the third overheads may be combined with make-up isobutane and co-fed said first stream as feed to the oligomerization reaction; and/or optionally the fractionator for separating C_4 's from the oligomers contains a bed of hydrogenation catalyst and hydrogen is fed to the fractionator such that dienes contained within said effluent are hydrogenated simultaneously with the separation. Fresh isobutane $_{30}$ may be fed to said second alkylation Zone.

FIG. 1 illustrates the overall process as it relates to the alkylation of an FCC C_4 steam with isobutane. As shown, isobutane enters the process via stream 101 and the FCC C_4 's enter the process via stream 102 . The FCC C_4 's are processed 35 prior to this step to remove any oligomerization catalyst poisons. Typically this can be accomplished by using a water wash column (not shown). The make up isobutane used for production of alkylate enters in stream 101 at a high purity, 1.e. $>$ 85 vol. $\%$. If such a stream is not available a more dilute, $\frac{40}{2}$ paraffinic, isobutane containing stream may be brought into the deisobutanizer, 23, for fractionation (not shown). The overall combined make up isobutane stream produced via the combination of streams 102 and 101 is depicted as stream 104. Stream 104, the rich isobutane stream, is then split into 45 two streams, 105 and 106. Stream 105 is fed to a second alkylation reactor as indicated below. Stream 106 is mixed with the FCC C_4 stream. The reason for mixing stream 106 with the FCC C_4 feed stream is for use as a heat sink to handle a portion of the heat of reaction during the oligomerization $\frac{50}{30}$ step. For water-cooled oligomerization reactors it is not nec essarily required and is thus optional.

The FCC C_4 in stream 102 stream and the optional isobutane rich stream in stream 106 are combined as stream 107 which is fed to the oligomerization reactor **18** containing a 55 bed 12 of oligomerization catalyst. The isobutene in the FCC C_4 stream is oligomerized in reactor 18 to form primarily diisobutylenes and triisobutylenes along with some co-dimers leaving a C_4 cut rich in normal butenes. The effluent from the reactor **18** in stream **108** comprising mixed C_4 60 and oligomers is fed to distillation column 19 wherein the oligomers are separated from the unreacted C_4 's. If desired a bed 22 of hydrogenation catalyst may be utilized in conjunc tion with a hydrogen feed via stream 115 to simultaneously provide removal of the dienes in the C_4 stream. The removal of the dienes helps the downstream alkylation units as diene presence increases the alkylation unit catalyst consumption. 65

The C_4 's (normal butene, and isobutane), are removed as overheads, condensed and removed via stream 109 which are later fed to a first alkylation system 22. Unreacted hydrogen is vented via stream 124.

10 15 The bottoms product oligomers from reactive distillation column 19 exit in stream 110 and are mixed with isobutane rich stream 105 to form a mixed alkylation feed stream 111 which is fed to a second alkylation system 20 wherein a portion of the isobutane reacts with olefins in stream 111 to form alkylate product. The effluent from the alkylation sys tem 20 exits as stream 112 and contains considerable isobu tane along with heavier alkylate products ranging from C_5 - C_{16} . Stream 112 is sent to a stripping or distillation column 21 in which the C_4 's are removed as overheads and product alkylate #2 is removed as stream 114. The C_4 's are removed as overheads via stream 113 are condensed and combined with streams 109 and 125 to form mixed stream 127 which is fed to the first alkylation unit 24 for production of alkylate by reaction with the olefins (butenes) in stream 109.

The effluent from the alkylation system 22 is removed in stream 116 and fed to a traditional deisobutanizer where alkylate product is removed as bottoms instream 117. Normal butane is removed in side stream 126 and isobutane is removed as overheads as stream 103 which is recycled to the process.

The whole point of the process is to produce more high quality alkylate at a facility and prevent the necessity of additional deisobutanizer capacity, which is a considerably large fractionation tower requiring significant energy usage for separation. Instead, for facilities having more than one alkylation system, the need for additional deisobutanizer capacity is averted by the addition of (a) an oligomerization reactor 18, (b) a fractionation tower 19 for separation of the oligomers from unreacted C4's and (c) a fractionation tower 21 for separation of alkylate product from the second alkyla tion system 20. The trade off on a piece count (numbering three new pieces of equipment) is valuable in that the replace ment pieces are small in size. For instance the required distillation stages for fractionators 19 and 21 can number less than eight stages each, whereas an additional deisobutanizer column will likely require 50-80 stages. This can significantly reduce overall equipment cost as less total steel is required to obtain the same quantity of motor fuel alkylate production. The process can be utilized with any type of alkylation pro cess, solid acid alkylation, HF alkylation, sulfuric acid alkylation and combinations thereof.

Alkylation With a Mixed C_4/C_5 Olefin Feedstock
Where C_4 and C_5 feed streams are alkylated a staged alkylation system approach can be used which is similar to that shown in FIG. 1, which is characterized as a process for the alkylation of isobutane, C_4 and C_5 olefins comprising the steps of:

(a) feeding isobutane and a stream containing C_5 olefins to a first alkylation Zone wherein a portion of the isobutane reacts with C_5 olefins to produce a first alkylate stream containing a first alkylate and unreacted isobutane;

(b) feeding said first alkylate to a debutanizing Zone wherein the first alkylate is separated as a first bottoms from a first overheads containing the unreacted isobutane;

(c) feeding the first overheads and a stream containing C_4 olefins to a second alkylation Zone wherein isobutane reacts with the C_4 olefins to produce a second alkylate stream containing a second alkylate and unreacted isobutane,

(d) feeding the second alkylate stream to a deisobutanizing Zone wherein the second alkylate is separated as a second

bottoms from an overheads containing the unreacted isobu tane. Preferably any normal butane contained in said isobu tane stream passes through said first and second alkylation systems and is removed as a side stream from said deisobu tanizer. Fresh isobutane may be fed to said second alkylation system to make up for that which is consumed as part of the reaction, and a portion of said first overheads may be recycled

Referring now to FIG. 2 alkylation feed consisting of FCC C_5 's with trace amounts of n-butane (<1 vol. %) can be used wherein an interim debutanizer 250 is utilized between two alkylation systems 230 and 240 which are processing C_5 's and C_4 's respectively. It is assumed that actual separation of the C_4 and C_5 olefins (not shown) occurs upstream of FIG. 2. The C_5 feed is sent to the first alkylation system 230 via 15 stream 201. The feed is mixed with the combined make up isobutane from stream 209 and the overheads from the deisobutanizer in stream 215 before being fed to the first system 230 as stream 202.

The first alkylation system 230 is used to produce a dilute 20 alkylate stream 204 containing a significant portion of isobu tane as the primary constituent. The effluent in stream 204 is sent to debutanizer 250 which produces a bottoms alkylate stream 205. The debutanizer 250 can be a stripper or a full distillation column requiring reflux. The debutanizer is used 25 instead of a traditional deisobutanizer to reduce the require ment of distillation stages and potentially the quantity of reflux and associated energy. The difference herein is that as the requirements are reduced away from a deisobutanizer operation, Small quantities of isopentane, residing in the 30 debutanizer overhead product are allowed to enter the second alkylation system 240. These minor quantities have a very minor effect on the overall alkylate quality obtained in alky lation system 240. Details of the effect are outlined below.

The resulting overhead stream 206 (containing primarily 35 isobutane, a minor amount of n-butane, a small amount of isopentane and possible containing trace amounts of 2.4dimethyl hexane and 2,2,4-trimethyl pentane) is fed along with either FCC C_4 's or an MTBE raffinate (containing mixed butylenes) in stream 207 to a second alkylation system 240 40 where they are allowed to react to produce a dilute alkylate product stream 208 in which a primary effluent constituent is isobutane. Stream 208 is then sent to a traditional deisobuta nizer 260 for separation wherein an isobutane rich overhead product is removed as stream 215 and recycled to alkylation 45 system. Alkylate product is removed as bottoms as stream 210. A normal butane rich stream is removed stream 212.

A significant variation of isobutane to olefin ratios in the two alkylation system and significant variability between alkylation for C_4 olefins versus alkylation from C_5 olefins can \sim 50 beachieved by utilizing flow lines 211 and 212. As one wishes to produce more alkylate from C_4 olefins flow line 212 is utilized. To product more alkylate product from C_5 olefins flow line 211 is used. The use of these flow lines stems from the alkylation system requirement to operate within a certain 55 isobutane to olefin volumetric ratio, varying from as low as 4:1 to as high as 15:1. As quantities of alkylate production from individual C_4 and C_5 feeds vary (i.e. total volumes of C_4 and C_5 olefins vary), the need for higher volumes of isobutane for the different alkylation systems becomes necessary. 60

In a manner similar to the use of flow lines 211 and 212, a modification of the process shown in FIG. 1 may be made which adds an additional isobutane rich stream 125 to serve as a bypass around the oligomerization section (reactor 18 and column 19). This allows for better energy utilization as it 65 reduces the duty on the distillation column 19. The inclusion of the bypass 125 depends on the overall isobutane to olefin

ratio desired in the alkylation systems and the exit require ments (temperature, pressure, number phases) set for the oligomerization reactor 18 as discussed earlier.

The ability to produce two separate alkylate products from stream 205 may be desired. The alkylate product in stream 205 may be fractionated in column 270. The alkylate product in overhead stream 213 contains higher octane and Reid Vapor Pressure (RVP) than that in bottoms stream 214. For certain regions using C_5 feed stocks, this becomes a necessary step in order to produce a primary alkylate meeting RVP specifications.

Alkylation With a Mixed C_3/C_4 Olefin Feedstock

For C_3 olefin feedstocks the same process flow scheme as in FIG. 2 may be used. Therein C_3 olefin would substitute in place of the C_5 olefin feed. Propane in the C_3 feed would be taken out in a depropanizer (not shown) residing in the first alkylation system 230.

Alkylation With a Mixed $C_3/C_4/C_5$ Olefin Feedstock

Combinations of the basic process shown in FIGS. 1 and 2 may be utilized for alkylation of a range of C_3-C_4 , C_4-C_5 and C_3 -C_s, olefin containing streams. FIG. 3 provides one basic option utilizing a single debutanizer 340 for alkylate effluent streams coming from olefin C_3 and C_5 feeds which have only trace amounts of normal butane in them. The C_3 olefin stream 301 is fed to alkylation system 320 while the C_5 olefin stream 301 is fed to second alkylation system 330. Make up isobu tane is fed as stream 303 and recycled isobutane from deisobutanizer 360 is added as stream 314 to make combined isobutane stream 304. Isobutane is provided to first alkylation system320 in stream 306 and to second alkylation system 330 in Stream 305. As noted both effluents 308 and 309 are fed to a single debutanizer 340 with the overhead in stream 310 being fed to a third alkylation system 340 which utilized FCC C_4 's from stream 307 to alkylate the normal butenes in stream 310. A bottoms alkylate stream is taken from debutanizer column 340 as stream 311 and deisobutanizer 360 as stream 312.

Along these lines more involved process schemes can be derived. For instance, the C_4 and C_5 FCC feed stream may be oligomerized separately. This would allow for more flexibil ity in how to alkylate the olefin streams which contain only trace amounts of normal butane. Additionally the step of oligomerizing the C_5 stream may be deleted with the C_5 FCC feed being fed directly to an alkylation system and then on to the debutanizer operation. Finally a single oligomerization unit may be used for both the C_4 's and C_5 's.

A process for the alkylation of isobutane C_3 , C_4 and C_5 olefins is characterized as comprising the steps of

(a) feeding a first stream containing isobutane to first and second alkylation zones;

(b) feeding a second stream containing propylene to the first alkylation Zone wherein the propylene reacts with a portion of the isobutane to produce a alkylate stream contain ing a first alkylate and unreacted isobutane;

(c) feeding a third stream containing C_5 olefins to the second alkylation zone wherein C_5 olefins react with isobutane to produce a second alkylate stream containing a second alky late and unreacted isobutane;

(d) feeding the first and second alkylate to a debutanizing Zone wherein the first and second alkylates are separated as a first bottoms from unreacted propylene and unreacted isobu tane as a first overheads;

(e) feeding the first overheads and a fourth stream contain ing C_4 olefins to a third alkylation zone wherein the C_4 olefins react with a portion of the unreacted isobutane in the first

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overheads to produce a third alkylate stream containing a third alkylate and unreacted isobutane;

(f) feeding the third alkylate stream to a deisobutanizing Zone wherein the third alkylate is separated as a second bot toms from the unreacted isobutane as a second overheads. Preferably any normal butane contained in said isobutane stream passes through said first, second and third alkylation systems and is removed as a side stream from said deisobu tanizer and the isobutane contained within said second over heads is recycled to said first and second alkylation systems 10

Staging and Energy

From a distillation perspective the invention is basic. Reducing the overhead specification to include minor amount of C_5 materials loosens the overall design requirements as compared to a traditional deisobutanizer, allowing for a sub- 20 stantial decrease in distillation column stages and optionally some reduction in column diameter, and condensing duty. The use of an inter-stage debutanizer between two alkylation units to reduce the need for additional deisobutanizer capac ity when expanding a C_4 MTBE raffinate alkylation operation to a full FCC C₄ operation, a C₄ alkylation operation into a C₃ and C_4 operation, a C_4 operation into a C_4 and C_5 operations or a C₄ operation into a C₃/C₄/C₅ operation, has not been heretofore described in the prior art. In the end, some amount of deisobutanizer capacity is required for the entire alkylation process flow scheme so that trace amounts of normal butane found in various feed streams (isobutane make up, olefin feeds, etc.) are not allowed to build up in the system. The invention reduces the overall deisobutanizer requirement necessary as compared with the traditional use of multiple deisobutanizers as depicted in U.S. Pat. No. 5,648,586. 25 35

Fundamental calculations can be made for comparison of a traditional expansion using two deisobutanizers for C_4 and C_5 alkylation (see U.S. Pat. No. 5,648,586) versus an expansion as disclosed herein as $F1G. 2.1$ he two cases can be compared $_{45}$ by simple distillation calculations for the effluent from the alkylation system which produces a dilute alkylate stream. A typical dilute alkylate stream is used for comparing the new scheme versus the traditional scheme for C_5 olefin feed containing isopentane and n-pentane. The dilute alkylate product composition as produced from an FCC C_5 feedstock and a recycle isobutane stream, and a dilute alkylate product pro duced form alkylating a C_3 olefin feed and isobutylene feed is provided in TABLE I. 55

Information presented by Kranz & Albright, "Alkylation of Isobutane with Pentenes Using Sulfuric Acid as a Catalyst: Chemistry and Reaction Mechanisms'. Ind. Chem. Res. 1992, 31, pp. 475-481, was used as a basis for the C_3 and isobutylene (iC₄=) cases. Although isobutylene is present herein it presents the basis used to represent the diisobutylene case shown in FIG. 1. US2004/017901 teaches that due to the de-polymerization behavior of diisobutylene, oligomers of 65 isobutylene act essentially like isobutylene on a weight basis in an alkylation system. 60

TABLE II provides the results of the comparison between the present interim debutanizer operation and a typical, prior art deisobutanizer operation for the three major feed cases: C_5 =, C_3 = and i C_4 =. The basis for these cases was to produce an alkylate product having less than a 5 psia RVP maximum. Distillation overhead pressure was set to 100 psia and the overall design was chosen to be at a factor of 1.2 times the minimum reflux requirement. The feed to the columns entered as a liquid at 100°F.

For each feed case two interim options are listed, each having a different control isopentane allowed in the overhead product. The "interim' stripper provides the lowest energy and staging option to meet the minimum RVP case of 5 psia, and thus allows a variable quantity of isopentane (present in the feed) to come out in the overhead product. The "interim' debutanizer case uses a maximum overhead isopentane speci fication of 0.25 wt% and limits C_4 's in the alkylate product to meet at minimum of a 5 psia RVP. It also requires an associ ated higher quantity of duty and staging to meet these speci fications when compared to the stripper case.

To an extent, the "interim' cases provide the range of operation in which an interim debutanizer operation may be configured for each of the feeds given in TABLE I. As feed composition changes variations of course exist. For the most part TABLE II provides a good view of the possible options and thus the basis for selecting the appropriate process for such a flow scheme for a C_5 , C_3 and isobutylene feed.

In terms of distillation staging required there is no com parison between traditional deisobutanizer requirements and the "interim' alkylation staging operations. Staging for all interim cases is always less due to the key components chosen for separation. Energy requirements for the "interim' cases with the present debutanizer operations typically require slightly higher duty as compared to traditional well designed deisobutanizer cases and with slightly lower duties required for the stripper cases. TABLE III provides a list comparing the relative staging required and relative energy require for the various "interim' options comparing the traditional debuta nizer, interim stripper and interim debutanizer.

DEBUTANIZER/STRIPPER/DEISOBUTANIZER COMPARISON								
	Reboiler Duty (BTU/Hr)	Minimum Reflux Ratio	Minimum Stages	Actual Reflux Ratio	Actual Stages	Alkylate RVP(psia)	Isopentane Wt % in OH	nC5 Wt5 in OH
			C_5 = Feed case					
Deisobutanizer	3.48E+07	0.47	33	0.57	84	To debut and depent	0.00	0.00
Interim C4 Stripper	3.07E+07			0	10	5	11.30	1.00
Interim debutanizer	4.18E+07	0.76	15	0.90	35	to depent	0.25	0.00
			C_3 = Feed Case					
Deisobutanizer	3.90E+07	0.47	33	0.57	84	To debut	0.00	
Interim C4 Stripper	3.30E+07			0	10	2.7	0.72	
Interim debutanizer	$4.04E + 07$	0.44	11	0.53	28	3.0	0.25	
			iC_4 = Feed Case					
Deisobutanizer	3.97E+07	0.47	33	0.57	84	to debut	0.00	
Interim C4 Stripper	3.35E+07			Ω	10	2.8	1.30	
Interim debutanizer	4.57E+07	0.65	13	0.78	31	3	0.25	

TABLE II

TABLE III

From just an energy and staging requirement, the stripper case is always preferred with a choice between traditional deisobutanizer and interim debutanizer becoming a clear 45 tradeoff between energy and staging and thus an operational cost versus initial capital cost consideration. A significant difference is that the interim designs allow some introduction of isopentane into a secondary alkylation system.

It has been found that the addition of isopentane to an $50₁$ alkylation system using $FCCC₄$ feed can cause reduced product octane if not properly controlled. As isopentane is limited to a number as low as 0.25 Wt % the difference in octane numbers between feed with and without isopentane becomes immeasurable due to the available precision of typical octane 55 tests. This allows better comparison of the options of tradi tional deisobutanizer with interim debutanizer on nearly equivalent terms, with the exception that the interim case provides for production of an alkylate product meeting RVP requirement. 60

It is anticipated that (1) with the use of more low pressure steam available for heating, (2) the use of available process streams for feed heating, and (3) the new retro fit options that are allowed due to the reduced staging requirements, a likely choice for a refiner between an interim debutanizer operation 65 and a traditional deisobutanizer will be that of the interim debutanizer as a means to increase their overall alkylation

capacity. For the case of C_3 olefin feed as shown in TABLE III $_{25}$ the interim options are clearly beneficial, with the stripper case having 12% of the staging, 85% of the energy and only 0.73 wt % isopentane in the overhead product due to the limited make of isopentane in the first alkylation system.

For the comparison made, the deisobutanizer case does not 30 include the total duty required for production of an alkylate product, whereas for all interim cases, except the debutanizer C_5 case, the associated duty provided allows for production of an alkylate product. For the C_5 case, the feed contains so much isopentane that the debutanizer bottoms stream 35 requires the use of a depentanizer to meet the alkylate RVP requirement. Thus additional equipment is required for that case. This is also true for the traditional deisobutanizer case.

Effect of Isopentane on a Secondary Alkylation System

As pointed out above, the fundamental effect of isopentane on product quality needs to be quantified. This was accom plished by performing an experiment wherein a mixture of isobutane and isopentane was co-fed with an FCC C_4 olefin to produce an associated alkylate product. The results are shown in FIGS. 4 and 5. As shown in FIG.4, increased isopentane in the feed causes a drop in alkylate quality as measured by the true (research+motor octane)/2 number as produced at a con stant olefin space velocity, total isoparaffin/olefin ratio, temperature and mixing energy. Comparing an overhead stream in the range of around 8% iC₅ in the iC₅-iC₄ mixture to one with none, only a slight drop in quality is seen, between 0.2-0.3 octane points. Within this range such quality effects are slight in comparison to the quality effects associated with
the overall alkylation system operation (OSV, I:O etc). Also it can be found from FIG. 5 that the cause for the reduction in octane up and (to some extent) beyond 8% is due rather to nificant conversion of isopentane during the alkylation process. This slight drop in quality is also offset in the scheme shown herein because isopentane, which is transferred to the next stage of alkylation (possibly reducing the overall octane of the second alkylate product) can consequently provide for the lower RVP and higher quality alkylate product in the first alkylation stage, depending on the base alkylate blending properties.

A particular finding, during the measurement of the effect of isopentane on product quality (FIG. 4), is that the net

isopentane consumption (using an FCC C4 feedstock) does not occur until the iC_s/(iC_s+iC₄) weight ratio is beyond 8 wt %. Indeed, contrary to a range of conditions listed in earlier studies (U.S. Pat. No. 5,583,275) it has been found that sul furic acid catalyzed alkylation (run at low temperatures of from 25-35° F., with acid strengths between 92-98 wt %) the incorporation of isopentane into alkylate is much reduced relative to isobutane, thus preventing overall consumption at the feed conditions. It was found and plotted in FIG. 4 that an $iC_5/(iC_5+iC_4)$ ratio of ≤ 0.1 provides for a net yield or net 10 make of isopentane when using an FCC C_4 feedstock which is directly opposite to that disclosed in U.S. Pat. No. 5,583,275.

The invention claimed is:

1. A process for the alkylation of isobutane with an olefin comprising: 15

- (a) feeding a first stream comprising an FCC hydrocarbon stream comprising an olefin to an oligomerization reac tor, wherein at least a portion of the olefins react with 20 each other to produce an effluent containing olefin oli gomers:
- (b) fractionally distilling the effluent from the oligomer-
ization reactor wherein unreacted C_4 olefins and lighter materials are separated as a first overheads from C_5 and heavier oligomers as a first bottoms;
- (c) feeding the first bottoms and a second stream contain ing isobutane to a first alkylation system where a portion of the isobutane is reacted with a portion of the olefin oligomers to form an effluent comprising a dilute alky late stream; 30
- (d) feeding the effluent from the first alkylation system to a debutanizer where C_4 's are removed as a second overheads and a first alkylate product is removed as a second bottoms;
- (e) feeding the first overheads and the second overheads to a second alkylation system where isobutane is reacted with C_4 olefins in said first overheads and said second overheads to form a second alkylate product.

2. The process according to claim 1 wherein said first 40 stream comprises an FCC C_4 stream containing normal butene, isobutene, normal butane and isobutane.

3. The process according to claim 1 where said first stream contains a mixture of C_4 and C_5 olefins.

4. The process according to claim 1 wherein said first 45 stream contains a mixture of C_3 and C_4 olefins.

5. The process according to claim 1 wherein said first stream contains a mixture of C_3 , C_4 and C_5 olefins.

6. The process of claim 1, further comprising feeding a third stream containing isobutane to the second alkylation system. 50

7. The process of claim 1, further comprising

(f) feeding the second alkylate stream to a deisobutanizing Zone wherein isobutane is removed as a third overheads, alkylate is removed as a third bottoms and normal butane is removed as a side stream. 55

8. The process of claim 1, further comprising feeding at least one of an FCC C3 and an FCC C5 stream to the first alkylation system.

9. A process for the alkylation of isobutane with butenes contained in an FCC C_4 stream comprising the steps of:

(a) feeding a first stream comprising an FCC C_4 stream containing normal butenes and isobutenes to an oligo merization reaction wherein the isobutenes react with 65 each other to produce an effluent containing C_5 and higher oligomers and normal butenes;

- (b) fractionally distilling the effluent from the oligomer ization reaction wherein the C_4 and lighter material is separated as a first overheads from the C_5 and heavier material as a first bottoms;
(c) feeding the first bottoms and isobutane to a first alky-
- lation zone to produce a first alkylate stream containing alkylate and unreacted C_4 's;
- (d) feeding the first alkylate stream to a debutanizing Zone wherein the unreacted C_4 's are removed as a second overheads and alkylate product is removed as a second bottoms;
- (e) feeding the first overheads and isobutane to a second alkylation Zone wherein the olefins in the first overheads are reacted with isobutane to produce a second alkylate stream containing alkylate, unreacted isobutane and unreacted normal butane;
- (f) feeding the second alkylate stream to a deisobutanizing Zone wherein isobutane is removed as a third overheads, alkylate is removed as a third bottoms and normal butane is removed as a side stream.

10. The process according to claim 9 wherein isobutane is fed as a second stream to the oligomerization reaction.

11. The process according to claim 10 wherein the third overheads is combined with said second stream and fed to the oligomerization reaction.

12. The process according to claim 9 where dienes con tained within said effluent are hydrogenated simultaneously with the fractional distillation in a bed of hydrogenation cata lyst with hydrogen.

13. The process according to claim 9 wherein fresh isobu tane is fed to said second alkylation Zone.

14. The process of claim 9, further comprising feeding the second overheads to the second alkylation Zone.

35 olefins comprising the steps of 15. A process for the alkylation of isobutane with C_4 and C_5

- (a) feeding a stream containing C_5 olefins to an oligomerization reactor, wherein the olefins react with each other to produce an effluent containing C_5 's and heavier oligomers:
- (b) feeding at least a portion of the oligomerization reactor effluent and isobutane to a first alkylation Zone wherein a portion of the isobutane reacts with C_5 oligomers to produce a first alkylate stream containing a first alkylate and unreacted isobutane;
- (c) feeding said first alkylate to a debutanizing Zone wherein the first alkylate is separated as a first bottoms from a first overheads containing the unreacted isobu tane;
- (d) feeding the first overheads and a stream containing C_4 olefins to a second alkylation Zone wherein isobutane reacts with the C_4 olefins to produce a second alkylate stream containing a second alkylate and unreacted isobutene;
- (e) feeding the second alkylate stream to a second debuta nizing Zone wherein the second alkylate is separated as a second-bottoms from a second overheads containing the unreacted isobutane.

60 first and second alkylation Zones and is removed as a side 16. The process according to claim 15 wherein any normal butane contained in said isobutane stream passes through said stream from said deisobutanizing Zone.

17. The process according to claim 15 wherein said first bottoms containing said first alkylate is fed to a fractional distillation wherein the first alkylate is separated into a third and fourth alkylate.

18. The process according to claim 15 wherein fresh isobu tane is fed to said second alkylation Zone.

19. The process according to claim 15 wherein a portion of said first overheads is recycled to said first alkylation Zone.

20. The process according to claim 15, further comprising feeding a stream comprising C_3 olefins to at least one of the $\overline{}$ first alkylation Zone and the second alkylation Zone.

21. A process for the alkylation of isobutane with C_3 , C_4 and C_5 olefins comprising the steps of:

- (a) feeding a first stream containing C_4 olefins to an oligomerization reactor, wherein the olefins react with each 10 other to produce an effluent containing oligomers;
- (b) feeding isobutane and a second stream containing at least one of propylene and isopentenes to a first alkyla tion Zone wherein the propylene, isopentenes, or both reacts with a portion of the isobutane to produce an alkylate stream containing a first alkylate and unreacted isobutane;
- (c) feeding the first alkylate to a debutanizing Zone wherein the first alkylate is separated as a first bottoms from $_{20}$ unreacted propylene and unreacted isobutane as a first overheads:
- (d) feeding the first overheads and at least a portion of the oligomerization reactor effluent to a second alkylation Zone wherein the oligomers react with a portion of the unreacted isobutane in the first overheads to produce a second alkylate stream containing a second alkylate and unreacted isobutane;
- (e) feeding the second alkylate stream to a deisobutanizing zone wherein the second alkylate is separated as a second bottoms from the unreacted isobutane as a second overheads.

22. The process according to claim 21 wherein any normal butane contained in said isobutane stream passes through said first, second and third alkylation Zones and is removed as a side stream from said deisobutanizing Zone.

23. The process according to claim 21 wherein the isobu tane contained within said second overheads is recycled to said first and second alkylation Zones.

24. The process according to claim 21, further comprising oligomerizing at least one of the C_3 and C_5 olefins prior to feeding to the first alkylation zones.

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