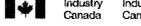


Industrie Canada

Canadä





Strategis Index:





Canadian Patents Database

12/20/2001 - 16:04:16 (12) Patent: (11) CA 526874

(54) HYDROCARBON SYNTHESIS

(71)		
(71)	(72) - 111 i (Country):	ISIDOR KIRSHENBAUM (Not Available) KENNETH K. KEARBY (Not Available)
(74) (45) June 26, 1956 (22) This case (43) Fig. (65) (52) Fig. (65) (51) Fig. (65) N/A This case are all cases No. No. 30) Letter the equation No. No. No. No.	(73) (Country):	ESSO RESEARCH AND ENGINEERING COMPANY (United States
June 26, 1956 (22) 1 5	(71) <u>i i d</u> (Country):	
(22) The conference (43) The conference (52) The conference of the	(74)	
(43) 1 2 2 252/8t (52) 2 3 3 3 3 4 4 5 4 5 4 5 4 5 4 5 4 5 4 5 4	(45)	June 26, 1956
(52) From State 1 and 1	(22) ** **	
(51) Brown to Longith N/A The transfer of the transfer of the No	(43)	
No Solve the restriction of the None None None None None None None None	(52)	252/81
None N/A Unknown	(51) The state that the last of the t	N/A
N/A Unknown	the state of the s	No
Unknown	(30) — portroins or production and the late	None
		N/A
	grand the second of the second	
	** Note: Data on abstracts and claim	ms is shown in the official language in which it was submitted.
** Note: Data on abstracts and claims is shown in the official language in which it was submitted.	t di sakarinin and a sa	
	/icw or Download Images :	
** Note: Data on abstracts and claims is shown in the official language in which it was submitted.		
	Cover Page Image	
** Note: Data on abstracts and claims is shown in the official language in which it was submitted. /iew or Download Images:	Abstract Image	
** Note: Data on abstracts and claims is shown in the official language in which it was submitted. /icw or Download Images: Cover Page Image	Claims Image	
** Note: Data on abstracts and claims is shown in the official language in which it was submitted. View or Download Images: Cover Page Image Abstract Image	Oisclosures Image	
** Note: Data on abstracts and claims is shown in the official language in which it was submitted. View or Download Images: Cover Page Image Abstract Image Claims Image	_	

The present invention rel tes to catalytic conversions and improved catalysts therefor. Note particularly, the invention is concerned with improved iron catalysts for the catalytic synthesis of normally liquid hydrocarbons and oxygenated compounds from CO and H₂.

Iron type catalysts are normally employed in the synthesis of hydrocarbons at relatively high temperatures of about 450°-800°F, and relatively high pressures of about 3-100 atmospheres abs. or higher, to obtain predominantly unsaturated and oxygenated products from which motor fuels with high octane ratines may be recovered.

The extreme temperature sensitivity and relatively rapid catelyst deactivation of the hydrocarbon synthesis have led, in recent years, to various attempts and proposals to employ the so-valled fluid catalyst technique wherein the synthesis gas is contected with a dense turbulent bed of finely divided catalyst fluidized by the gaseous reactants and products. This technique permits catalyst replacement without interruption of the process and greatly improved temperature control. However, the adaptation of the hydrocarbon synthesis to the fluid catalyst technique has encountered serious difficulties, particularly when iron catalysts are used.

Application of the fluid technique requires case of fluidization and attrition resistance in addition to the conventional characteristics determining catalyst activity, such as total desired wield and active catalyst life. It is also desirable that the catalyst be active in the temperature range above 600°F, and still be highly selective to C_h + hydrocarbons, since under these conditions high octane motor fuels are obtained. None of the prior art iron catalysts complies satisfactorily with all of these requirements.

Iron catalysts are usually prepared by the reduction of various natural or synthetic iron oxides or by the decomposition of iron carbonyls,

the catalytic activity being enhanced by the addition of such promoters as various compounds of alkali metals or the oxides of chromium, zinc, aluminum, megnesium, manganese, the rare earth metals and others in small amounts of about 1-10%. While some of these catalysts exhibit excellent activity characteristics they are without exception deficient with respect to ease of fluidization, and/or attrition resistence particularly when used in commercial runs of several hundred hours duration. Even fluidized catalysts obtained from sintered iron, which have been found to exhibit excellent fluidization and attrition characteristics show signs of disintegration in long run operation.

of mechanical strength during operation has been found to be closely connected to a high rate of cerbon deposition on the catalyst, encountered at the conditions required by the synthesis using iron catalysts. The catalyst disintegration which accompanies excessive carbon deposition is believed to be the result of a migration of carbon into the iron lattice by the mechanism of interstitial carbide formation followed by disintegration of the carbide to free carbon. This process may continue until the catalyst mass contains about 90% of carbon.

It will be appreciated from the above that ar iron catalyst of satirfactory synthesizing activity, selectivity, and catalyst life which may be used in commercial operation without substantial catalyst disintegration and carbon deposition is a need strongly felt in the synthesis art. This drawback of iron catalysts has been the major obstacle in all attempts to apply the fluid catalyst technique to the iron-catalyzed hydrocarbon synthesis. The present invention overcomes this obstacle,

It is, therefore, the principal object of the present invention to provide improved iron catelysts for the catalytic synthesis of hydrocarbons from CO and $\rm H_2$.

A further object of this invention is to provide an improved hydrocarbon synthesis process operating in the presence of iron catalysts

which are not subjected to excessive disintegration and carbon deposition.

A more specific object of this invention is to provide an improved hydrocarbon synthesis process employing the fluid catalyst technique
in the presence of iron catalysts of highest disintegration resistance
throughout runs of commercial length.

Other and further objects and advantages of this invention will appear hereinafter.

In accordance with the present invention, carbon deposition on iron synthesis catalysts is substantially reduced and catalyst disintegration correspondingly suppressed while activity, selectivity and catalyst life are maintained at highest levels, when the iron preferably in the form of its carbonate is combined with a metal carbonate base which in itself exhibits a high disintagration resistance and the composite is subjected to a reduction treatment. While a wide variety of metal carbonates may form, in combination with iron, catalysts of good activity, selectivity and disintegration resistance, superior results have been consistently obtained by combining iron carbonate, particularly ferrous carbonate, with the carbonates of metals of Group II of the Pariodic System, and more particularly of one or more of the metals of calcium, barium and zino. The relative proportions of the elements in the unreduced catalysts of the present invention may very within wide limits. However, this iron content should not be substantially lover than about 0.53 by waight or about 1% by weight of ferrous carbonate, with the other metal carbonate being present in major proportions. Active carbonate-supported catelysts with over-all iron concentrations of less than 0.5% by weight may also be made provided that the iron is properly distributed on the surface of the catalysts. A distribution of about 5-40% by weight of ferrous carbonate and about 60-90% by weight of the carbonate of a Group II metal is generally preferred.

An outstanding and surprising characteristic of the carbonate catalysts of the present invention, particularly those comprising berium and/or calcium carbonate, resides in the fact that they efford excellent

conversion and liquid product yields even in the absence of the conventional catalyst promoters such as halides, carbonates or orides of alkali metals which prior to the present invention have been considered indispensable for catalysts of satisfactory activity and selectivity.

While it is not wished to limit this invention to any specific theory or probably reaction mechanism, it is believed that the carbonate bases give the catalysts the proper alkalinity required for high selectivity to liquid product.

Examples of catalysts useful for the purposes of the present invention include composites consisting of 90% BaCO3 and 20% FeCO3; 80% ZnCO3 and 20% FeCO3; 80% ZnCO3 and 20% FeCO3; etc.

These catalysts may be prepared by any suitable method of coprecipitation, precipitation of the iron component on the precipitated
support, impregnation or mechanical mixing, known per se in the art of
catalyst manufacture, followed by reduction. A simple and expedient
method of adding the iron, especially in very low concentrations, to the
surface of the catalyst is by the decomposition of iron carbonyl upon the
catalyst. Although a number of different iron compounds such as the
oxides, oxalates, acetates, etc. may be used; superior results have been
obtained when using the method of precipitating ferrows carbonate on the
precipitated metal carbonate support.

The invention will be further illustrated by the following specific examples.

BRAUPLE I

About 1141 gms. of BaCO₃, made by the addition of amnonium carbonate to a solution of FaCl₂ in distilled water, was slurried in 3 liters of water. A solution of 344 gms. of FaCl₂,4H₂O in 2 liters of distilled water was slowly added to this slurry. Contact with air was kept at a minimum during these steps. The slurry was stirred for one hour and an additional 149 gms. of FaCl₂,4H₂O dissolved in 600 c.c. of water was added. Then 285 gms. of amnonium carbonate, dissolved in 1 liter of water,

were stirred in and the mixture was further stirred for one hour. The precipitate was permitted to settle, washed free of chloride by decantation, and the resulting sludge was dried over-night in an atmosphere of CO₂ at about 300°F, and then pilled. The catalyst contained about 20% by weight of RaCO₃ and 20% by weight of FeCO₃. It was reduced prior to use with about 1000 volumes of H₂ per volume of catalyst per hour for 4 hours at about 900°F.

The catalyst so prepared was tested in a fixed bed laboratory unit at about 630°-650°F., about 250 lbs. per sq. in. pressure, a throughput of about 200 volumes of synthesis has per volume of catalyst per hour, and an W2:00 ratio of about 1:1. The results are summarized below:

CC Conversion, \$\beta\$ 96
Ch+ Yield, c.c./cu.m.
H2+CO Consumed 211

Carbon Formation
Selectivity to C,Vol.\$\beta\$
CC to \$\beta\$ 0.97

Selectivity to 0,% of Reference* 32

Catalyst Age
Cu. ft. of CO Converted
per 1b. of Catalyst
69
% CON After Use
22 (Agreement within ex% CON Theoretical for
Bacon
27

The above data show that carbon formation on this catalyst is only a fraction of that observed on the reference catalyst, while liquid product yields at high temperatures are excellent in spite of the absence of any catalyst promoter. However, low carbon formation is also evidence of a lower disintegration tendency for catalysts having a comparable abrasion resistance.

The activity and selectivity of this catalyst are substantially maintained even after high temperature calcination. A sample of the fresh

 Basis of comparison is the carbon formation on a reduced catalyst consisting of 99% precipitated iron oxide and 1% potassium fluoride. catalyst was calcined for about 4 hours at 1600°F, before reduction at 900°F. The catalyst so prepared was tested in a fixed bed laboratory unit at about 630°-650°F, about 250 lbs. per sq. in, pressure, a throughput of about 200 volumes of synthesis gas per volume of catalyst per hour, and an H₂:00 ratio of 1:1. The results are summarized below:

CO Conversion, \$\\ C4+\ Yieli (c.c./cu,m. H2+CO Cons)\ C3+\ \ "\ \ "\ \ "\ \ "	94 182 237
Carbon Formation Selectivity to C, Vol. 5 CC to C Selectivity to C, 5, of Reference*	0.71 44
Cetalyst Age Cu. ft. of CO Converted mer lb. of Catalyst	7.44

EXAMPLE II

A catalyst was prepared as described in Example I. However, 20.2 gms. of $K_2^{\circ}CO_3$ was stirred into the washed paste prior to drying. The precipitate paste was then dried and pilled as described in Example I to form a catalyst consisting of 80 parts by weight of BaCO₃, 20 parts by weight of FeCO₃ and 2 parts by weight of $K_2^{\circ}CO_3$. The catalyst was reduced and tested at the conditions of Example I. The results were as follows:

Co Conversion, % Cu+ Yield, c.c. per cu.m. H2+CO Consumed	96 211
Carbon Formation Selectivity to C. Vol. 3 CO to C Selectivity to C. 3 of Reference	1.9
Catalyst Age Cu. ft. CO Converted per 15, of Catalyst	126

The data of this example show that the addition of a promoter to the carbonate catalyst of the present invention has no appreciable influence on liquid yields or carbon formation.

^{*)} Basis of comparison is the carbon formation on a reduced catalyst consisting of 99% precipitated iron oxide and 1% potassium fluoride.

EXAMPLE III

A solution of 344 gms. of FeCl₂.4H₂O in 1 liter of water was slowly added to a slurry of 800 gms, of CnCO₃ in 4 liters of water. To this mixture there was then added slowly with stirring a solution of 225 gms. of ammonium carbonate in 1 liter of water. After stirring for 1 hour, the precipitate was permitted to settle and was washed free of chloride ion by decantation. The precipitate was then dried at 250°F, in a stream of CO₂ and pelleted.

The catelyst was reduced at 900°F, and tested under the conditions of Example I with the following results:

●CC Conversion, 5		96
*CO Conversion, 3 Cu+ Yield (cc/M3	Cons.)	205

The second of th

Catalyst Age
Cu. ft. CO Converted
per 1b. of Catalyst
% CON After Use
% CON Theoretical for CaCON
Supported Iron Catalyst
49

Carbon Formation
Selectivity to C, Vol. % CO to C 0.40
Selectivity to C, % of Reference 16

EXAUPLE IV

The catalyst was prepared as described in Example III. However, 2002 gms, of $K_2^{CO}_3$ was stirred into the washed precipitate pasts. The precipitate was then further treated as described in Example III, the final catalyst containing 80 parts by weight of $CaCO_3$, 20 parts by weight of FeCO₃ and 2 parts by weight of $K_2^{CO}_3$. A test was carried out at the conditions of Example III and had the following results:

CO Conversion, %	86
Ch+ Yield, c.c./cu. m, of	
H ₂ +CO Consumed	200+

Carbon Formation
Selectivity to C, Vol. \$ 00 to C 0.40
Selectivity to C, \$ of Reference 23

Catalyst Age
Cu. ft, of CO Converted per
lb, of Catalyst 133 .

The data of Examples III and IV show that the carbonate supported catalyst of the invention effords high activity and high liquid product sensitivity at high temperatures and gives low carbon formation, even in the absence of conventional catalyst promoters.

Control of the Contro

...

Y SIGHAKE

A solution was made containing 733 cms. of zinc chloride and 275 cms. of ferrous chloride (YeO₂,4H₂O) in 2 liters of distilled water. An emmonium carbonate solution was made by dissolving 1480 cms. of armonium carbonate in 6 liters of water. The two solutions were then added simultaneously with stirring to 5 liters of distilled water. After stirring for 1 hour, the precipitate was washed free of chloride ion and dried at 250°F. Contact with air was kept at a minimum during these steps.

This catalyst was reduced at 900°F, and tested in a fixed bed laboratory unit at about 600°F, about 250 lbs. per sq. in. pressure, a throughput of about 400 volumes of synthesis gas per volume of catalyst per hour, and an H₂:00 ratio of about 1:1. Under these conditions, a 00 conversion of 95% was obtained. The C4+ yield (cc. per cu. m. of H₂+00 consumed) was 160. These data show that Group II metal carbonates other than barium and calcium are suitable as supports for highly active and selective iron catalysts even in the absence of conventional catalyst promoters.

EXAUPLE VI

The catalyst was prepared as described in Example V. However, 135 gms. of the filter cake was slurried with water and 2.7 mms. of $K_2^{00}_3$ was then stirred into the paste. This paste was dried and treated as described in Example V, the final catalyst containing °0 parts by weight of $Z_1^{00}_3$. 20 parts by weight of $Z_1^{00}_3$ and 2 parts by weight of $Z_2^{00}_3$.

This catalyst was tested at 650°F, but otherwise at the conditions of Example V. Under these conditions a 60 conversion of 96% was obtained. The C_h^+ yield (c.c. per cu. m. of H_2^+ CO consumed) was 200+.

These results show that the zinc carbonate supported catalyst is more sensitive to the addition of promoters than the barium and calcium

carbonate catelysts. However, both types of cetalysts exhibit a promoter effect clearly different from that of other types of iron catelysts. This is demonstrated by the data of Example VII below.

In this example it is seen that both $\text{Li}_20.\text{Fe}_{23}^0$ type catalyst and a zinc oxide promoted catalyst require promotion with K co in order to give high selectivity to C_h+ .

EXALPLE VII

A catalyst having the composition Li₂0.Fe 0 was made by mixing 444 gms, of lithium hydroxide into a paste of 942 gms, red iron oxide in 200 c.c. of distilled water. After ball-milling for 15 hours and drying over-night at 240°F, the catalyst was calcined at 1600°F, for 3 hours.

A promoted catalyst having the composition by weigh' of 99
Li₂0.Fe₂0₃-1K₂00₃ was prepared by mixing 5 gms. K₂00₃ into a paste containing 643 gms. of the dried Li₂0.Fe₂0₃ poste made as above. After drying at 240°F., the catalyst was calcined at 1600°F. for 3 hours.

A zinc oxide based ion type catalyst was prepared by adding slowly 1016 ems. of Fe(NO₃)₃.9H₂O dissolved in 2 litera of distilled water to a slurry of 1106 ems. ZnO in 3 liters of water. The slurry was stirred until all of the iron had precipitated. The precipitate was filtered, washed, reslurried, filtered and washed. Half of the filter cake was then dried at 250°F, and calcined for 3 hours at 250°F. The final catalyst contained 80 parts by weight of ZnO and 20 parts by weight of Fe₂O₂.

The other half of the washed filter cake was made into a thick paste and impregnated with 5 cms. $K_2\text{CO}_3$. The paste was dried and then calcined for 3 hours at \$50°F., the final composition being about \$0 parts by weight of ZnO, 20 parts by weight of Fe $_2\text{O}_3$ and 1 part by weight of $K_2\text{CO}_3$.

These catalysts were reduced at 900°F, and tested at temperatures above 600°F, with a feed gas containing an $\rm H_2/CO$ ratio of about 1. The results are summarized below:

with $1/1~\mathrm{H}_2$ and CO feed. The results are tabulated below:

Catalyst	FeCO3	(Synthetic)	Siderite
Promoter	None	2% K CO	None
Temperature, °F.	600	600650	600
Feed Rate, v/v/hr.	400	400	200
CO Conversion, % CO+H2	96	94	92
Feed Rate, v/v/hr. CO Conversion, % CO+H2 Ch+ Yield, (c.c./cu. m. H2+CO Cons.)	173	162	100

In the foregoing examples the use of a single metal carbonate as the catalyst support has been shown. It will be understood however, that similar results may be obtained when using suitable mixtures of the metal carbonates disclosed.

While the above experimental data were obtained in fixed bed operation, the relative comparisons hold for fluid operation, even though the higher gas throughputs, high recycle ratios and high catalyst turbulence typical for fluid operation quite generally cause a slight decrease of conversion and liquid product yields and an appreciable increase of carbon formation and catalyst disintegration. It follows that the catalysts of the invention, as a result of their greatly reduced carbonization and disintegration tendency and their superior liquid product selectivities even at high temperatures coupled with long catalyst life, are perticularly useful for fluid catalyst operation and in this respect greatly superior to other catalysts of the iron type. Catalysts, in accordance with the invention, suitable for fluid operation may be prepared substantially as outlined in the above examples and sized to particle sizes of about 20-150 microns, preferably 50-100 microns. The conditions of fluid synthesis operation are well known in the art and need not be specified here in any great detail for a proper understanding of the invention by those skilled in the art. Briefly, these conditions may include catalyst particle sizes of 20-200 microns, superficial linear cas velocities of about 0.1-3 ft,/sec., bed densities of about 10-120 lbs. per cu. ft., H_2 :00 ratios of about 0.5-3, gas recycle ratios of about 0-5, temperatures of about 550°-750°F.,. and pressures of about 150-650 lbs. per sq. in.

Catalyst Base	Li ₂ 0.Fe ₂ 0 ₃		30 ZnO-20 Fe ₂ O ₃	
Promoter	None	1% K2co3	Mone	1% K ₂ 00 ₃
Temperature, °F.	650	650	650	550
Fressure, psig.	250	250	250	250
Feed Cate, v/v/hr.	400	300	200	200
CO Conversion, 3	. 90	96	95	97
Ch+Yield (cc/cu.m.	•	•	,,,	,,
CO+H2 Cone.)	100	172	108	196
Hours on Stream	246	54	207	209
Selectivity to C.				•
(% of Reference)	80	105	10	2 _β

The above data illustrate the entremely strong effect of alkali metal promoters on liquid product and carbon selectivities, which is typical for usual iron type synthesizing catalysts. It will be observed that the catalysts of the invention differ widely in this respect from non-carbonate catalysts of the iron type.

It has also been found that the metal carbonate supported iron catalysts of the invention are far superior to catalysts consisting essentially of iron carbonate and this in spite of the much higher iron content of the latter. This is demonstrated by the following further example.

EXALPLE VIII

Ferrous carbonate was precipitated by the simultaneous slow addition of aqueous solutions of ferrous chloride and ammonium carbonate to water under stirring. The precipitate was allowed to settle and washed free of chloride by decentation. One-half of the washed precipitate was dried in an atmosphere of SC and pilled.

The other half of the washed precipitate was mixed with K CO in 2 3 amounts sufficient to incorporate about 2% of K_2^{CO} in the final catalyst, dried and pilled as described above.

Natural Siderite, containing FeCO₃ (in a concentration corresponding to 40.4% Fe) and with impurities of magnesium, silicon, manganese, calcium and aluminum was crushed to a powder, calcined at 1000°F, in a closed pan, pilled and recolcined at 1000°F.

These three carbonate catalysts were reduced in hydrogen and tested at temperatures of 600°F, and above, and at a pressure of 250 psig.

The present invention is not to be limited by any theory of the mechanism of the process or catalyst nor to any exemples given merely for illustration purposes, but only by the following claims in which it is wished to claim all novelty inherent in the invention.

- 1. An improved iron catalyst for the synthesis of hydrocarbons from carbon monoxide and hydrogen, consisting essentially of a reduced composite of a major proportion of the carbonate of at least one metal selected from the group consisting of calcium and barium amounting to not substantially less than about 60% by weight and a minor proportion of iron amounting to not substantially less than about 0.5% by weight.
- 2. The catalyst of Claim 1 in which said iron is present in said composite prior to reduction in the form of a carbonate.
- 3. The catalyst of Claim 2 in which said carbonate is ferrous carbonate.
- 4. The catalyst of Claim 1 which contains a small amount of an alkali metal promoter.
- 5. The catalyst of Claim 4 in which said promoter is potassium carbonate.
- 6. The catalyst of Claim 1 which has a fluidizable particle size of about 20-150 microns.
- 7. The catalyst of Claim 1 in which said iron is present in said composite prior to reduction in the form of ferrous carbonate, said major proportions are about 60-95% by weight and said minor proportions about 5-40% by weight.
- 8. The catalyst of claim 7 in which said major proportions are about 80% by weight and said minor proportions about 20% by weight.

 $\psi_{/}$