Special Real L27
Kaiser Wilhelm Institute Researches on Synthetic Fuels
Translation by Max leva
TOM REEL 101

Research on Fe and Fe-Cu Catalysts

Performance under Varying Synthesis Conditions

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## The Lehavior of Iron Contacts During the Operation with Hydrogen-rich Synthesis Cas (Fe-Contact + 1/h% K2 CO3)

Fore recent experiments in which iron catalysts were used in benzine synthesis have shown that those catalysts will give yields which are very nearly equal to those obtained when using cobalt catalysts.—Hydrocarbons obtained with iron-catalysts—compare favorably also to those obtained by cobalt when one compares their anti-knock properties. The catalysts must be used at temperatures of around 235-260°C., that is, approximately 45-70° higher than cobalt catalysts. Then carrying out the middle-pressure synthesis with cobalt catalysts, we have made use of an apparatus which was cooled by steam. It was our desire to use the same set-up for experiments with iron catalysts, without having to make any experimental changes. It would have been advantageous therefore to be able to work at a lower operating temperature. This succeeds when one uses a gas which is richer in hydrogen than a normal gas used along with the iron catalyst.

A time test with mixed gas showed that it is possible to start the synthesis at 210°C. After 15 days of operation and working at 213°C., a yield of 100 g./Nm<sup>3</sup> of ideal cas was still obtained, and the increase in temperature to assure an economical yield was only small. After operating for 180 days, the necessary operating temperature to assure a feasible yield was 235°C.

On the tenth day of the synthesis, 97% of the CO gas was used up, whereas 50% of the initial N2 constituent remained unconverted. In order to keep the yields up, it was necessary to operate in two stages, and the gas leaving the first stage and entering the second stage had to be replonished with sufficient CO to bring its final composition up to its original value. Pefore replenishing the gas however, for other reasons) it was desired to see what would happen if the gas issuing from stage one was conducted into stage two.

It may be seen that thus it was possible after 70 days of operation to obtain a yield of 157 g./ $N_m^3$  of ideal gas when operating in two stayes.

In order to lower the temperatures still more, experiments were carried out in several (4) stages. A gas of composition CO:Ho = 1:4 was used.

In Table II, the results of such a run may be observed. For the first hit days, the second stage was operated with an initial gas having the composition of the end gas from stage #1, including the gasol hidrocarbons. After the hith day, the second stage was operated with a gas having been replenished with 00; however, the gasol had been removed. The third and fourth stages were operated with a gas as was obtained from the previous stages and the gasol was left in.

We were able to demonstrate that iron catalysts may be used at temperatures below 200°C., giving economical yields and producing only little CO<sub>2</sub>. Generally, it was observed that at the lower temperatures, more water formed than CO<sub>2</sub>. The CO conserption is satisfactory, and after the third stage, approximately 90% has been used up. The fourth stage was operated at 170°C, and after 20 days of operation, it was still not necessary to raise the temperature. The CO conversion amounted to 60%,

The yields of stages one and two, were tabulated and referred to  $1\,N_{\rm H}^{-3}$  of initial gas as fed to the first stage (III). It was found that after the second stage,  $106\,\rm g_{\circ}$  of hydrocarbons/ $N_{\rm H}^{-3}$  of ideal gas were obtained. It was furthermore discovered that the gasol in the gas passing over the catalyst in stages 3 and 4 was converted into higher hydrocarbons and no longer could be quantitatively determined as gasol.

Average analysis of exit gas from Stage 1.

CO2	4.3%
8KW	
CO	7.4
H <sub>2</sub>	76.4
KW	3.8
N <sub>2</sub>	7.0

On adding 11.7% CO to this gas, one obtains the following gas:

3.9%
1.0
17.1
68.4
3.4
6.2

If the second gas has to be compressed into a 40-liter bomb up to a pressure of 160 atmospheres, one requires:

•	•
CO2	6.39
8KW	1.6
CO	27.3
H <sub>2</sub>	109.3
KW	5.5
N2	710.0

Wheinpreussen-Gasol was available with 32% skW and 68% kW; therefore only 1.6 atm. skW had to be added. At the same time, 3.4 atm. kW were compressed into the bomb; however, since 5.5 atm. kW were required, it was necessary to add an additional 2.1 atm. of CH<sub>1</sub>. The mixing of the gases was easily accomplished therefore in this manner.

In order to refer the individual yields of the 4 stages to one cubic meter of the original starting gas, it is permissible to work in the individual stages with gases containing gasol. During those experiments, however, the conditions were kept as nearly as possible to actual conditions because it appeared undesirable from the point of view of experience that the gasol of the end-gas should be removed after each stage. The yield determinations of the third and fourth stages show, however, that it is possible to obtain economic quantities of benzine when using iron catalysts (IV).

#### Recycling Experiment

Recycling in teneral gives similar results to those obtained when working in various stages. Therefore, a recycling experiment was undertaken with a furnace containing 18 reaction tubes and filled with 180 g. of catalyst.

The activity of the catalyst was checked with a CO-rich gas at 180°C., and then operation was started with a gas containing 1CO + 1112. The operating temperature was 180°C. The conversion of CO was checked for both recycling as well as one single passage through the tubes. The work was carried out with 24 to 30 liters of end-gas, and approximately 100 liters were circulated. Consequently, during the recirculation, a 3 to 4 times higher flow velocity was obtained in the reaction furnace.

	1830	1880	2030
	Circulated circulated	Mot Circulated circulated	Not Circulated circulated
-Contraction	26% 27%	33.5% 28%	31.5% 21.6%
C-balance	42.5 g./Nm3 32.5 g./im3	31.2 g/Nm <sup>3</sup> 30.6 g./Nm <sup>3</sup>	31.2 Pa/Nm3 211.7 g/Nm3
CO conversion	49% 47.5%	The second of the control of the con	61.5% 52%

The preceding table shows contractions, carbon balance, and CO conversion at three different temperatures. From this experiment, the A.K. benzine was removed at 193°C. (without recirculation), and a distillation was carried through of the constituents boiling at 200°C. The boiling point indicated approximately normal pentans, normal hexame, normal heptane, and normal octane. A yield determination for the experiments of the 18-tube furnace at 193°C. and 30 liters of end-gas per hour showed 41.5 g./Nm³ of ideal gas. This yield was obtained without recycling of the end-gas. No comparative yield was obtained in the recycling experiment because the recycling pump failed to operate after-some time.

#### Flow Experiment

The influence of the flow velocity was studied for the same catalyst at different temperatures (10 g. of iron or 16 g. of iron catalyst were charged into the furnace). The following table shows the end-gas quantities, the total gas quantity converted, and the converted CO per hour at various temperatures.

-Temporature	Liters end-gas per hr.	Percent con- traction	Liters A.G. per hr.	Converted gas quantity per hr.	con- CO
180°.	2	37	3.20	1,200 1	0,456 1 80
	4	20	5.00	1,000 1	0,390 1 44
	8	17	9.65	1,650 1	0,660 1 39
190	2	28	2.78	0,780 1	0,430 1 69
	4	21	5.26	1,260 1	0,544 1 56
	8	11	9.30	1,300 1	0,600 1 33
200°	2	35	3.08	1,080 1	0,531 1 90
	4	27	5.48	1,480 1	0,635 1 61
	8	20	10.00	2,000 1	0,830 1 址
of works printing a local				esservene po grane. Par	

From the tabulation, it is observed how for every temperature the contraction and the CO conversion reduced (about 50%) as the end-gas quantity increases from 2 to 8 liters per hour. Nevertheless, the total quantity of gas and the total quantity of CO converted increases steadily, so, for instance, at 200°C., approximately twice as much gas is converted for 8 liters of end-gas than for 2 liters of end-gas. The yield at 190°C. for 8 liters of end-gas was 31.1 g. of hydrocarbons per normal cubic meter of ideal gas.

# The Influence of Different Modes of Catalyst Induction Upon the Operation of Hydrogen-Rich Starting Cas

The catalysts have been protreated in the already well-known manner. The either worked with L liters of pure CO (per 10 g. of iron) for 25 hours (100 liters altogether) at 325°C. and 1/10 atm.; or we operated for 2-1/2 hours with LO liters of CO per hour per 10 grams of iron (100 liters altogether) at 325°C. and 1/10 atm.

In order to investigate the influence of different pretreatments of the catalyst upon operation with hydrogen-rich gas, the first thing we did was that we treated a catalyst according to the earlier methods long enough with mixed gas at 220°C.—255°C. and atmospheric pressure until it gave a contraction of 30% under the same conditions (4 liters per 10 g. of iron per hour). Then we increased the pressure to 15 atm. and a gas of 100 + 4H<sub>2</sub> was used at 180°C. and 2 liters per hour of endgas. It was found that the yield even after a temperature of 200°C. was reached after 6 days of operation, was not as high as the yield when the catalyst was pretreated under reduced pressure.

	Temperature_	Percent Contraction	CO conversion, percent	Days of operation
After pretreatment	1810	12	20	1
with mixed gas	189	<b>1</b> 2	25	5
	196	22	37	6
	200	20	ili	7
	199	20	-46	11
	200	20	32	177
	199	~ 20	34	20
	199	17	31	21
After pretreatment	180	30	71	1
in vacuum with pure	co 181	2U	60	5
• • • • • •	180	24	66	_8
	180	21	58.5	10
	179	16.5	38.5	<b>18</b> :
	187	23	48.5	26
	190	34	52	30
	190	23	50	43

when the two methods of pretrentment are compared with each other, it may be noticed that the vacuum treatment has the advantage. With the catalyst having been subjected to the vacuum treatment, the temperature had to be raised to 190°C. only after operating for as long as 30 days. Also, the time of operation, the amount of carbon monoxide converted was higher than in the case of the catalyst pretreated under ordinary pressures. Since in both cases, the same catalyst was

used (28), it is out of the question that the differences in yields could be explained with a loss of activity of the catalyst. An experiment was carried out with a catalyst pretreated in vacuum and operated under normal pressure with a hydrogen-rich gas (1:1). The results are tabulated in the next table and compared to those of the normal protreated catalyst of the same constituents.

	Pretreated		1:1	Not pretreated					
Tempera ture	7. Table 1.	con-	No. of days	Temper- ature	Con- trac- tion	CO con- version	lio. of days		
181	4. *		1	180	6		<b>351</b>		
185	7		5	180	5		:2		
195	-6	18	3	187	- 5		2		
219	10	21	5	200	- 5	***	$\mathbf{Z}$		
240	12.5	47	. 10	<b>210</b>	7	12	- 8		
21.0	15	51	11	210	7	23	- <b>9</b> -		
540	19	115	15	209	6	20	13		
****				230	16	1:0	15		
				<del></del> .			· · · · · · · · · · · · · · · · · · ·		

It was not possible to obtain any appreciable amounts of hydrocarbons below 200°C. Then working at 240°C, or above, the conversions became normal. Approximately the same results were obtained in both cases when a hydrogen-rich gas was used under normal pressure and low temperatures.

Another experiment was carried out with the same catalyst. However, it was pretreated. The operation was carried through at 15 atm. with hydrogen-rich starting gas (1:1). The operating temperature was kept at 190°C. at first; however, the CO conversion under those conditions was only 10%. When the temperature was raised to 235°C., the contraction increased to 15%, and the CO conversion went up to 38%.

## Pressure Series

The influence of operating pressure upon the middle-pressure synthesis using iron catalysts was investigated at 0, 1.5, 3.5, 7.5, 15, and 30 atms. The following table shows the effect of the various pressures. In all cases, the catalyst used was the same as described before, and was pretreated prior to all experiments in a vacuum by using pure CO.

O atm。 (s	eo above te	ible)	.1.5 atm.				3.5 atm.				
Con tra Temp. tion	c- CO-con-	No. of days	Temp.	Con+ trac- tion		No. of days	Топра	Con- trac- tion	Co con-	No. of	
181 li 185 7 195 6 219 10 210 12 210 15 210 19	51.	1 2 3 5 10 11 15	180 180 180 189 190 190	9 17 15 26 22 8 6	29 43 27 58 52 23 31	2 7 10 21 27 37 39	181 180 181 180 180 180 189	33 37 20 20 11 23 31 26	60 91 60 57 18 12 59 58	2 7 10 16 23 32 37 47	

	7.5 obn.			15 ntm.					30 atm			
agmor	con- trac- tion_	GO con- version	lo.ol ayab	Tomp.	Con. trac- tion	co con= version	No. of days	Temp.	con- trac- tion	CO con- vorsion		
181 180 180 181 182 180 -180	34 38 25 30 26 32 27 21	52 70 71 68 68 55	10 15 22 32 36 10	180 181 180 180 179 179 187 190	30 21 21 17 19 23 -34 23	60 66 58 38 33 48 62 50	15 8 10 18 24 26 30 43	180 175 178 180 179 180 179 190 190	*60 50 33 30 25 19 18 20 31	100 100 75 56 61 39 39 42 53 65	2 7 4 5 0 0 1 1 2 2 5 1	

The experiments at O atm. and 1.5 atm. are unsatisfactory at low temperatures. The run at 3.5 atm. showed a 40% conversion of CO after a month of operation (at 180°C.). The experiment at 7.5 atm. showed the best results. After 50 days of operation at 180°C., the conversion had not decreased below 50%, and up to this time only a slow decrease in activity of the catalyst was observed. The yield determination between the 34th and 38th day of operation still showed 40.5 g. of total hydrocarbons per Nati of ideal gas. "hon working under 15 atm., it was necessary to raise the temporature from 180 to 190°C. after 25 days of operation. A yield determination between the 16th and 20th days of operation showed 34.5 g. of total hydrocarbons per Mm3 of ideal pas. The 30 atm. run showed an initial activity corresponding to 100% CO conversion. In order to avoid over-heating of the catalyst, the temperature had to be dropped to 175°C. However, it could be raised again to 180°C, on the 6th day, llowever, on the 20th day of operation, the conversion had dropped to 42%, and a temperature increase to 190°C. was unavoidable. However, this increase did not help indefinitely. The yield determination between the 9th and 13th days of operation showed 45 g. of total hydrocarbons per Nm of ideal gas. From all of these experiments, it may be observed that the most favorable pressure is 7.5 atm. when using a hydrogen-rich gas of composition 100 + 4H2. It is speculated that the best CO conversion occurs somewhere around 10 atm. (see tabulation 5)

#### Experiment 1:6

In order to lower the working temperature some more, when working with iron catalysts, an experiment was carried out with an initial gas of composition 100 + 6H2 (same catalyst and same pretreatment as before). The experiment was started at 15 atm. and 160°C.

Tenperature	Contraction	CO conversion	llo. of days
1500	31%	70%	ż
159	26	62	<b>.3</b>
158	29	62	Ĭ.
160	22	55	8
160	24	55	13
160	22	60	18
160	20	55	20
160	20	45	21

The carbon balance on the 13th to 21st days of operation showed 23 g. of condensable hydrocarbons per 1m3 of pas. It was remarkable how little CO2 was formed.

Only approximately Diff of the end-gas consisted of CO.

This experiment also proves that benzine may be produced at lower temporatures when using Iron catalysis than the temperatures of the middle-pressure synthesis using cobalt.

Concerning the products of this reaction, it may be said that the pararties obtained were of remarkably light color when using a hydrogen-rich gas. When a CO-rich gas was used, the products of the iron and middle-pressure syntheses were brown. The results have been given already on the distillation analyses of the fractions boiling below 200°C. (page 1)

## One Percent Alkali Experiment

An experiment was carried out at 15 atm. and 180°C., with the cutalyst containing one percent potassium carbonate. The cas used consisted of 1CO + 1H2. It was found that the activity of this cat-lyst was considerable lower. We obtained yellow oil without solid paraffin.

### Fe-Cu-Contact for Operation with Hydrogen-Rich Starting Cas-

Sarlior experiments shows that small additions of copper improved the activity of iron catalysts. In order to investigate this effect with a hydrogen-tich gas, Fe-Cu-Catalysts were compounded. The catalysts used contained 5 parts of Fe and one part of Cu. The copper was co-precipitated with the iron. After the precipitation, the precipitate was made alkaline with I/MS of potassium carbonate.

## Pressure Series with Fe-Cu Catalysts

The Ferri-Cu-catalyst (5:1) was pretreated in a vacuum and was used for three experiments. One experiment was run at 0 atm., another at 1.5 atm., and a third at 15 atm. pressure, using a hydrogen-rich starting gas. The following tabulation shows the results of the experiments.

O atm.				1.5 atm.				15 atma			
Temp.	Con- trac- tion	No. of	CO con- version	Temp.	Con- trac- tion	No. of	CO con- version			- No. of days	CO con- version
185 215	5 15 16	1 5 16 19 31	6 21 23 20	180° 180 180 190 190 193	14: 13: 24: 27: 13:	2 8 10 13 16 29	20 24 45 52 42	182 181 180 185	32 32 32 32 32 32 32 32 32	1 6 10 15 21 25 33	59% 46 56 61 59 18

A comparison with the corresponding experiments carried out with a copper-free catalyst shows that for 0 and 1.5 atm., no improvements were observed. At 15 atm., it appears that a small increase in catalyst activity was noticed (see also tabulation 6).

## Normal Pressure Experiments with Pe-Cu Catalysts

Another series of experiments were run with Fe and Cu catalysts and normal pressures and various starting gases containing different proportions of hydrogen to carbon monoxide. Both Ferri- and Ferro-copper catalysts were used at normal temperature, and were tested for their activity by using mixed gas; then catalysts of the same composition were used with hydrogen-rich gas at correspondingly lower temperatures

	1 : 2	Ferrige Gu - Contact					
		Andrew Mark State (1965) The Company of the State (1965)		l : 2 (Parallel-Test)			
Marth Village 1	Percent	CO con-	<u> </u>		<b>Percent</b>	Percent CO con-	
2,414 2 3 4 1 4 1 4 1	Contraction	version	Days	Temperature	Contraction	version	Days
<b>=235°</b>	21		2	2350	22	Control Control	2
236	32		3	235	25		4
229	26	80	4	233	27	- 2000	5
229	28	70	8	231	25		7
230	22		9	245	30		- 8
231	18		13	€ 7	F		
245	33	<b></b>	<u> 1</u> j	<u> </u>			
	1:4			1:6		transkir.	
201	13	24	2	200	6	-	2
201	18	(30)	3	200	·5		h
200	14	24	4	205	. <b>6</b>	-	5
205	10	27	6	207	୍ରି <b>3</b> ₁	•••	8
210	9	-	8		etys gradition		
100 miles (1990) 1800 miles (1990)		Ferr	o - Cu	- Contact			
1 	1:2			-1:2	(Parallel-Te	st)	
236	34	90	2	231	-31		2
232	30	-	3	228	30	90	3
232	32	90	Ĭ.	227	34		7
229	28	90	8	229	30	90	8
231	31	90	10	230	33	90	10
238	25		19	230	34		11
	1:4			1:6			
201	<u>l</u> i		2	205	<b>3</b> .		2
210	12	30	3	209	10	28	3
213	11	59	7	210	13	40	4-
215	13		10	210	ŭ	43	8
	te de Cara e car			215	17	70	10

The higher activity of the Ferro-Copper catalysts is especially apparent when operating with mixed gas at temperatures of above 230°C. But when working with the hydrogen-rich gases at around 200-215°C., this higher activity is scarcely noticeable.

When the experiments are compared with those employing a corpor-free not pretreated catalyst, no important improvement can be noticed over a copper-containing catalyst for the temperature range of 200-210°C