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TECHNICAL REPORT No. 110-45

WARTIME RESEARCH ON SYNTHETIC FUELS BY THE

KAISER WILHELM INSTITUT FUR KOHLENFORSCHUNG

June 1945

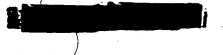
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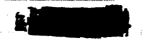
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# TECHNICAL REPORT No. 110-45

WARTIME RESEARCH ON SYNTHETIC FUELS BY THE KAISER WILHELM INSTITUT FUR KOHLENFORSCHUNG

#### SUMMARY

Wartime research by the Kaiser Wilhelm Institut fur Kohlenforschung on the Fischer-Tropsch synthesis has been primarily directed along the line of iso-paraffin synthesis. Good yields of iso-paraffins have been obtained using the normal synthesis gas at three hundred (300) atmosphere pressure and four hundred twenty (420) to four hundred fifty (450) degrees centigrade over an alumina-thoria catalyst. Zinc oxide-alumina has also been used successfully. About one hundred twenty (120) to one hundred thirty (130) grams per cubic meter of synthesis gas have been obtained as liquid products of over ninety (90) percent iso-paraffins.

Synthesis of aromatics has been studied, using thirty (30) atmosphere pressure and five hundred (500) degrees centigrade. However, very poor yields and napthene side reactions do not make this process look promising. Further work was a bandoned by KWI for the duration of the war.

Fischer-Tropsch synthesis using iron catalysts was studied, but inferior operations resulting from this catalyst made it only desirable as a wartime substitute for scarce chromium.

Juno 1945

U. S. NAVAL TECHNICAL MISSION IN EUROPE

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WARTIME RESEARCH ON SYNTHETIC FUELS BY THE KAISER WILHELM INSTITUT FUR KOHLENFORSCHUNG

### 1. DETAILS OF RESEARCH

# (a) <u>Iso-Paraffin Synthesis</u>

- (1) Operating conditions for the synthesis of isoparaffins have been found to lie in the range of three hundred (300)
  atmospheres pressure and four hundred twenty (420) to four hundred
  fifty (450) degrees centigrade. For pressure under three hundred
  (300) atmospheres the yield falls of f rather sharply, although a
  very slow reaction will take place at thirty (30) atmospheres or
  ever. Higher pressures than three hundred (300) atmospheres give
  increasingly greater percentages of exygenated products, until at
  one thousand (1000) atmospheres the principal product is dimethyl
  other. Similarly, lower operating temperatures give slower reaction
  rates, more unsaturated compounds, a very high percentage of alcohols,
  and less carbon formation on the catalyst. For temperatures slightly
  greater than four hundred fifty (450) degrees centigrade, the products
  become principally napthenic, and carbon deposition becomes excessive.
  Instantaneous reaction rates, however, are increased.
- (2) It has been found that the best synthesis gas has a CO/H<sub>2</sub> ration of 1.2 volumes of CO to one (1) of H<sub>2</sub>. Increasing the hydrogen increases methane formation; decreasing the hydrogen lowers the overall yield.
- an aluminathoria co-precipitated one, although ZnO-A<sub>1</sub>2<sub>0</sub>3 appears to be nearly as good and much cheaper. Using either of these materials it was found necessary to burn off the carbon deposits about every two weeks of steady operation. This was accomplished with air at the temperature of the synthesis (450 degree centigrade). Catalysts so treated have been used continuously for ever six (6) months without appreciable decline in activity. Their heat sensitivity is also quite good, as they have been held for prolonged periods at eight hundred (800) degrees centigrade without damage.

## 1. DETAILS OF RESEARCH (a) (cont'd).

- (4) Heat evolution during synthesis is approximately the same as with the normal synthesis, i.e. one-fifth (1/5) of the heat of combusion of the products. Since it is possible to work in a twenty (20) to thirty (30) degree centigrade temperature range, this lessons the problem of very close temperature control normally encountered in this process. Another advantage is found in the fact that sulphur does not seem to be nearly as deleterious as in former syntheses, although the upper allemable limit has not yet been determined.
- (5) Gas volocities have been fairly accurately studied, and it was determined that twenty (20) cubic continuotors of a two (2) to four (4) millimeter catalyst were necessary for each ten (10) liters of synthesis gas per hour. Any increase above this velocity gave sharp decreases in yield and also tended to form increasing amounts of alcohols.

(5) Product yields are as follows from a ZnO-Al<sub>2</sub>O<sub>3</sub> catalyst using a CO/H<sub>2</sub> ratio of 1, at three hundred (300) atmospheres and four hundred fifty (450) degrees contigrade:

and four hundred fifty (450) degroes contigrado:

Total yield of C<sub>3</sub> and higher 120 - 130 gm./m<sup>3</sup> 19 - 19.4 4 = 20 gm./m<sup>3</sup> 3.0 - 3.0 15

C<sub>4</sub> (90% isobutane) 50 - 80 gm./m<sup>3</sup> 1.5 - 11.9 15 × 1.50 = 20

C<sub>5</sub> and higher (over 97% iso) 30 - 60 gm./m<sup>3</sup> 4.5 9.0 22.4

(7) A comparison of the two (2) catalysts shows that for a thoria-alumina catalyst (1:4 by weight) the best yields of all me obtained. Higher gas velocities can be used, and there is little tendency to form alcohols. However, there is a greater tendency to form carbon on the catalyst, meaning shorter burn-off times. Experiments have varied the ratio of the two materials in the catalyst from twenty (20) to forty (40) percent Al<sub>2</sub>O<sub>3</sub>, with very little change in everall yields. However, increased alumina does increase methane formation somewhat. The addition of one-half (½) to one (1) percent K2OO<sub>3</sub> to the catalyst will give a slight increase in yield.

# 1. DETAILS OF RESEARCH (a) (cont'd).

- (8) On the other hand the AnO-Al<sub>2</sub>O<sub>3</sub> catalyst (1:1 by weight) is believed to carry more promise as a commercial catalyst because of its cheapness. It does give slightly less yield than the theria type, and produces about ten (10) percent alcohols. There is less tendency to form carbon deposits on the catalyst. The same tendency holds true as with theria catalysts, ic., higher alumina content promotes methane formation. However, experiments have been made over a range of Al<sub>2</sub>O<sub>3</sub> content from 2:1 to 1:2 without much change in results. Additions of K<sub>2</sub>CO<sub>3</sub> do not appear to give any advantage.
- (9) The theria type catalyst was prepared by using two (2) liters of therium and aluminum nitrates at their boiling point and in the right proportion to give a 1:4 weight ratio of theria to alumina. This was added one (1) liter of a boiling Na<sub>2</sub>CO<sub>3</sub> solution of proper strength to give one hundred (100) grams of dry co-pet., in stoiciometric amounts. Here dilute solutions gave greater density catalysts. The resultant precipitate was washed, dried, and sized to two (2) to four (4) millimeters.
- (10) The same method was used for proparing the ZnO type catalysts, except it was found that the addition of the nitrate solution to the soda solution instead of the normal method gave increased liquid hydrocarbon yields.
- (11) An alternate method for proparing theria catalysts, and the one which gave the best yields, was to make sodium aluminate and then precipitate with sulphuric acid. Therium hydroxide precipitate was also made, and these two ppts., were washed separately and mixed while wet. They were then air dried at one hundred (100) degrees centigrade and ground to normal size.

## (b) Aromatic Synthesis

(1) Aromatics were successfully synthesized by the use of Cr, No. The exides as catalysts. Five (5) to ten (10) percent K2CO3 was added to reduce carbon formation. It also reduced activity. A CO/H2 ration of 1:1 was used, at thirty (30) atmospheres and five hundred (500) degrees contigrade. Any higher temperature or pressure gave excessive carbon formation on the catalyst, while

### 1. DETAIL OF RESEARCH (b) (cont'd)

lower temperatures gave no aromatics, and reduced the reaction rate. Lower pressure similarly reduced the reaction rate.

- (2) The yields were very poor, being only about eight (8) to ten (10) gm./m of liquids which were about fifty (50) percent aromatics and fifty (50) percent napthenes. There was lots of methane formed, and much unconverted gas. The aromatics were principally toluene, xylene, and alkyl benzones. The napthones were a grand mixture not yet identified.
- (3) Because of the poor yields and the excessive carbon formation when working at such high temperatures, the Kill staff had temperarily abandoned this line of research to concentrate on more productive wartime work.

#### (c) Iron Catalyst Synthosis

- (1) The entire staff of the KWI was quite emphatic in stating that the use of iron catalysts in Fischer-Tropsch synthesis was considered only as a wartime necessity because of the shortage of chromium. Cobalt catalysts were stated to be definitely superior, cleaner to use, and capable of operating at lower temperatures. They did not know of a single instance where iron catalysts were actually used in commercial installations, although it had been planned in the event that chromium became unobtainable.
- (2) Iron catalysts are prepared by precipitation from its nitrate with soda solution, drying at one hundred ten (110) degrees contigrade, grinding and sizing, and then treatment with 1:1 CO/H<sub>2</sub> gas at reduced pressures (from 0.1 to 1 atmosphere) and two hundred forty (240) to three hundred twenty (320) degrees contigrade. This treatment forms an iron carbide on the catalyst surface, and is continued until a sharp drop in CO<sub>2</sub> content of the exhaust gas indicates cessation of carbide formation. It is generally used in sizes ranging two (2) to five (5) millimeters.

# 2. CONCLUSIONS AND RECOMMENDATIONS

The development of the Fischer-Tropsch process as a producer of iso-paraffins, and thus a source of high quality aviation and automotive fuels, opens the way for it to become the most versatile of all synthetic fuel processes. It would then be capable of producing products which range all the way from high quality diesel fuels and lubricants to excellent aviation gaseline. No other process could make such a claim. It is therefore recommended that this information be made available to all interested agencies, and that the work of Dr. Pichler and Dr. Koch of KVI, which is still continuing at the Institute, be closely followed in order to keep American technology as well informed as possible.

#### 3. DETAILS OF SHIPMENTS

catalysts used in iso-paraffin synthesis are being forwarded to Bureau of Ships (Code 341) on Consignment Tag No. 3655. Under the same consignment is also being shipped small samples of n-octane, n-nonane, and n-decame for use in the standardization of mass spectrographic analysis procedure.

Propored by:

D. R. DEWEY, Licut. (jg), USNR