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CATALYST FOR SYNTHETIC-METHANOL PRODUCTION.

No Drawing. Original application filed May 26, 1926, Serial No. 111,884. Divided and this application filed October 14, 1926. Serial No. 141,665.

gen, and pertains more directly to the preparation and employment of improved catalysts containing cobalt in the process.

Methanol may be produced by combining oxides of carbon with hydrogen in the presence of a suitable catalyst at elevated temperature and pressure. Carbon monoxide, carbon dioxide, and mixtures of the two oxides may be employed, these substances reacting with hydrogen according to the following equations:

Carbon monoxide— $CO + 2H_2 \rightleftharpoons CH_3OH$ Carbon dioxide— $CO_2 + 3H_2 \rightleftharpoons CH_3OH + H_2O$

It is observed that when carbon dioxide is the oxide employed, one molecule of water is formed for every molecule of methanol produced. On the other hand when pure carbon monoxide is used, theoretically there is nothing produced by the reaction but methanol. Actually in practice pure carbon monoxide and pure carbon dioxide are both difficult to obtain economically, so that the methanol synthesis is carried out by reacting a mixture of carbon monoxide and car-

bon dioxide with hydrogen.

In addition to the reactions producing methanol there are, in the methanol synthesis, undesirable side-reactions which cut down the yield of the desired product. The principal side reaction which may occur is the formation of methane, which is illustrated

below:

CO+3H₂≠CH₄+H₂O CO₂+4H₂≠CH₄+2H₂O

In addition to the methane side-reaction there are other side reactions which sometimes occur in which there are produced esters, aldehydes, organic acids, ketones, and hydrocarbons other than methane; these reactions occurring as the result of the poly-

merization or condensation of methanol or

its decomposition products.

When a gas mixture comprising carbon oxides mixed with an excess of hydrogen over the amount theoretically required to produce methanol is passed over a catalytic deleterious to the methanol reaction—namely substance comprising metals or their oxides the "methanating metals"—iron, nickel, or 105 at a pressure above 50 atmospheres and at a cobalt. In our improved catalyst compo-

Our invention relates to the production of temperature above 250° C. there is nearly methanol by the high pressure catalytic always produced some reaction between the 55 combination of oxides of carbon and hydrogaseous components. The extent of this reaction depends to some degree on space velocity, temperature, and pressure, but the fact remains that under the conditions outlined, carbon oxides and hydrogen react to 60 some extent in all cases.

The substances formed by such a process depend both as to identity and as to amount, almost entirely on the nature and activity of the catalytic substance present. The meth- 65 anol catalysts proposed in the past have been

of two principal types:—i. e.—

1. Mixtures of finely divided metals, or, what is equivalent, mixtures of easily reducible metal oxides;

2. Mixtures of oxides of metals non-reducible, or difficulty reducible under the conditions of the methanol synthesis.

In the past, also, one class of easily reducible metal oxides, that is the oxides of 76 iron, nickel, and cobalt have been described as absolutely worthless for the production of methanol catalysts since these metals under normal conditions prevent the formation of methanol, the reaction producing only 80 methane.

We have discovered a new type of methanol catalyst which produces a higher yield of methanol than those formerly employed in the art, and at the same time produces a 85 very pure methanol, uncontaminated by byproducts. An additional advantage accruing from the use of our catalysts is that the formation of methane in the process—with the consequent destruction of valuable hy- 90 drogen—is practically eliminated.

We have now discovered a new type of valuable methanol catalyst which comprises

three main elements-i. e.

(1) One or more non-reducible metal ox- 95 ides such as zinc, magnesium, cadmium, chromium, vanadium, tungsten, etc.; (2) One or more easily reducible metal

oxides such as copper, silver, iron, nickel, cobalt, etc.;

(3) A metallic halide.

It will be observed that in our improved catalyst we may employ substances normally 2 1,625,997

sitions these normally deleterious substances serve advantageously, apparently exerting only a normal hydrogenating catalytic action productive of methanol. Or we may 5 employ easily reducible oxides such as those of copper or silver which are already known in the art as advantageous components of certain methanol catalysts. In these cases however, our improved catalysts containing 10 halides produce far better results than those known in the art for otherwise identical catalyst mixtures.

The precise method by which the metallic halides produce the improvement is not 15 known to us. At least three theories may be used to account for the phenomenon-

namely:

(1) The metallic halide acts as an ordinary "promoter" in increasing catalyst ac-

20 tivity; (2) The metallic halide interacts with other metallic oxides present in the catalyst to produce oxychloride compounds which, in turn, either serve as catalyst promoters, or 25 may so modify the purely physical state of the mass as to produce a more active cata-

(3) The metallic halide may be reduced by the high pressure contact with hydrogen 30 and carbon oxides during the commencement of its use as a catalyst and the resultant finely divided metal may increase the catalyst activity, or alternatively, the metallic halide may first react with some other metal producing the halide thereof, which may be reduced in the same manner.

However, the precise explanation for the improved result attained by the addition of metallic halides to methanol catalysts is not known and forms no part of our invention.

To produce our improved catalysts it is not necessary that the amount of metallic halide added bear any exact weight relation to the remaining constituents of the catalyst, though the amount of halide added should preferably be less than one chemical equivalent of the amount of principal metallic oxide present in the catalyst.

As illustrative of our invention we will 50 cite two specific catalysts containing cobalt. In order to indicate the comparative efficacy of these catalysts in methanol production we have selected a standard set of conditions of use as follows. The reported yields of condensate from the methanol reaction by use of the following catalysts is based on the effect produced when a gas mixture comprising 8% carbon dioxide, 3% of carbon monoxide and 89% hydrogen is passed through 1 liter of catalyst granules at a space velocity of 75,000-100,000, at a pressure of 2000 pounds, and at a temperature of 340-400° C.

In selecting a standard set of conditions it is our intention only to thus display the catalyst efficacy and by no means to limit

the scope of our invention. Modification of the conditions of the reaction will, of course, modify the results, but the results will always be proportionate to the catalyst employed.

For example, an increased space velocity 70 produces an increased hourly yield and an increase in operating pressure has the same effect. Likewise the use of pure carbon monoxide as distinguished from carbon dioxide or a mixture of the two results in an 75 increased methanol percentage in the condensate, and, since carbon monoxide seems to react more readily, an increased condensate volume.

Example I.

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2250 grams of cobalt nitrate (Co(NO₃)₂.-6H₂O) are dissolved in 250 liters of water and the solution is heated to 95° C. There is then added 7.5 kilograms of zinc oxide 85 after which the solution is vigorously stirred and sufficient ammonium hydroxide is added to precipitate the cobalt as cobalt hydrate. The precipitated mass is recovered by decantation and filtration, is washed, dried, 90 and broken up into granules. The granules are then moistened with a solution containing 1480 grams zinc chloride and dried once

The hourly yield of condensate is about 95 1.5 liters and contains about 58% of methanol.

Example II.

If the zinc chloride in Example I is re- 109 placed by 1300 grams of magnesium bromide the hourly yield of condensate and percentage yield of methanol is not greatly modified.

This application is a division of our co- 105 pending application Serial No. 111,884, filed May 26th, 1926. In the said application we have claimed, generically, methanol catalysts comprising difficultly reducible metal oxides, easily reducible metal oxides, 110 and metallic halides and therefore such generic claims are not made herein.

Now having described our invention, we claim the following as new and novel.

1. A methanol catalyst initially compris- 115 ing a plurality of difficultly reducible metal oxides, cobalt oxide, and a metallic halide.

2. A methanol catalyst initially compris-

ing a difficultly reducible metal oxide, cobalt 120 oxide, and a metallic halide.

3. A methanol catalyst initially comprising zinc oxide, chromium oxide, cobalt oxide, and a metallic halide.

4. A methanol catalyst initially comprising zinc oxide, cobalt oxide, and a metallic 125 halide.

5. A methanol catalyst initially comprising zinc oxide, chromium oxide, cobalt oxide, and zinc chloride.

6. A methanol catalyst initially compris- 130

chloride.

7. A process for the production of synthetic methanol which comprises passing a mixture of hydrogen and carbon oxides at a temperature of 350-450° C. over a catalyst initially comprising a plurality of difficultly reducible metal oxides, cobalt oxide, and a metallic halide.

8. A process for the production of synthetic methanol which comprises passing a pressure in excess of 50 atmospheres, and at a temperature of 350-450° C. over a catalyst initially comprising zinc oxide, chromium oxide, cobalt oxide and a metallic halide.

9. A process for the production of synthetic methanol which comprises passing a pressure in excess of 50 atmospheres, and catalyst initially comprising zinc oxide, chromium oxide, cobalt oxide, and zinc 25 chloride.

In testimony whereof we affix our signatures.

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ing zinc oxide, cobalt oxide, and zinc catalyst initially comprising a difficultly chloride.

reducible metal oxide, cobalt oxide and a