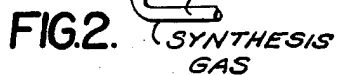


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2 Sheets-Sheet 1



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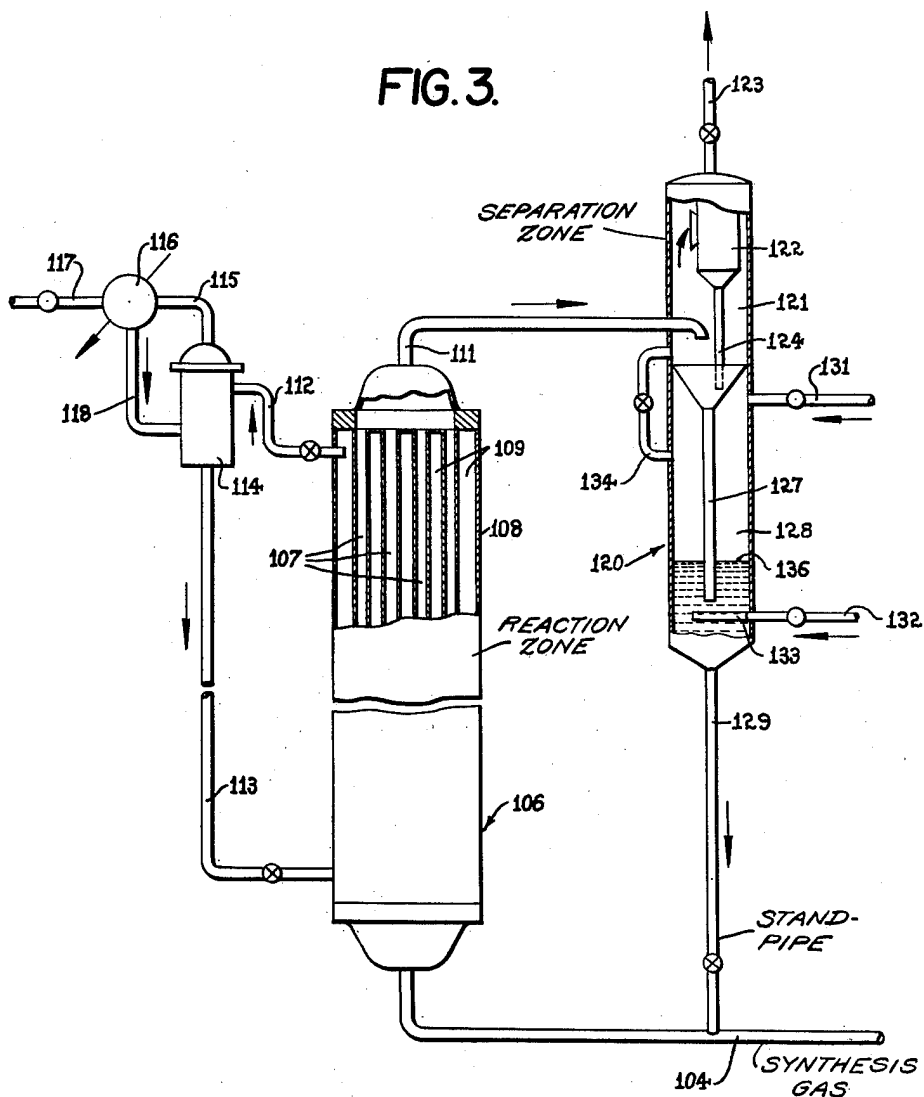
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HIGH VELOCITY PROCESS FOR SYNTHESIS OF ORGANIC COMPOUNDS

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2 Sheets-Sheet 2

FIG. 3.



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HIGH VELOCITY PROCESS FOR SYNTHESIS
OF ORGANIC COMPOUNDS

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9 Claims. (Cl. 260—449.6)

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This invention relates to the synthesis of organic compounds. In one aspect this invention relates to the hydrogenation of an oxide of carbon under conditions such that organic compounds having more than one carbon atom per molecule are produced. In another aspect this invention relates to a method and apparatus for the synthesis of organic compounds by the hydrogenation of carbon monoxide with a relatively low ratio of hydrogen to carbon monoxide in the presence of a finely-divided powdered catalyst.

It has been known for some time that hydrogen and carbon monoxide may be made to react exothermically in the presence of certain catalysts and under specific reaction conditions to form hydrocarbons and oxygenated organic compounds having more than one carbon atom per molecule. In general, the synthesis of hydrocarbons by the hydrogenation of carbon monoxide is accomplished in the presence of a metal or an oxide of a metal chosen from group VIII of the periodic table as a catalyst at pressures below about 500 pounds per square inch gage and at temperatures below about 750° F.

Various methods have been practiced to effect the reaction of hydrogen and carbon monoxide to produce organic compounds. Among these methods are those known as fixed-bed catalyst operations and fluid-bed catalyst operation. The fixed-bed operation comprises passing a reaction mixture of hydrogen and carbon monoxide through a stationary bed of catalyst in a reaction zone, and the fluid-bed operation comprises passing a reaction mixture through a finely-divided catalyst mass suspended in the reaction mixture in the reaction zone. Characteristically, certain reaction conditions are necessary for each of these processes and for the particular catalyst used.

The synthesis feed gas or reaction mixture comprises a mixture of about 1 to 2 mols. of hydrogen per mol of carbon monoxide and may be prepared by the catalytic conversion of natural gas, steam, and carbon dioxide.

The most recent development in the synthesis of organic compounds from hydrogen and carbon monoxide has been in the fluid-bed type operation. This type of operation has had several apparent advantages over the fixed-bed oper-

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ation and has yielded organic compounds of high quality and in large quantity. In such a fluid-bed operation at a temperature of about 600° F. and at super-atmospheric pressures using a fluidized iron catalyst, a contraction of about 41 per cent to about 70 per cent and a carbon monoxide disappearance of about 85 per cent to about 100 per cent have been observed. The selectivity of the reaction indicates that about 25 per cent to about 40 per cent of the CO is converted to CO₂, and oil and water yields of about 100 to 130 cc. per cubic meter and about 60 to 120 cc. per cubic meter, respectively, are obtainable.

Even in view of the relatively good results obtained by the fluid-bed type operation certain inherent disadvantages have been found: In such fluid-bed operations in which the catalyst is suspended in the reaction gas, classification of the catalyst often occurs. There is also a tendency for the fluid bed to settle after extended use of the catalyst due to wax and carbon accumulation on the catalyst. These tendencies have required certain design considerations in maintaining certain distances between the heat transfer surfaces. Furthermore, special consideration must be made for the removal of heat liberated in the reaction, and with fluid-bed operation specially constructed apparatus is necessary for the removal of such heat with attention directed to the fluidized catalyst itself. It is much to be desired, therefore, to provide a process and apparatus which overcome these difficulties, at least partially.

It is an object of this invention to provide a process for the synthesis of organic compounds having more than one carbon atom per molecule.

It is another object of this invention to produce hydrocarbons and oxygenated compounds by the reaction of carbon monoxide and hydrogen in the presence of a hydrogenation catalyst.

Another object of this invention is to provide an improvement in the synthesis of hydrocarbons in the presence of a finely-divided fluidized catalyst in the reaction mixture.

Still a further object is to provide a method for the synthesis of hydrocarbons using a relatively low feed ratio of hydrogen to carbon monoxide.

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Still another object is to provide a fluidized process for the hydrogenation of carbon monoxide in which the catalyst life is extended and prolonged.

Various other objects and advantages will become apparent to those skilled in the art from the accompanying description and disclosure.

Much to our surprise, we have found that an oxide of carbon may be hydrogenated to yield organic compounds having more than one carbon atom per molecule in the presence of a finely-divided hydrogenation catalyst which is present in the gaseous reaction mixture in a much smaller amount than heretofore thought possible. We have found that a catalyst concentration in the gaseous reaction mixture of less than about 18 pounds per cubic foot of gas is adequate to carry out the reaction between hydrogen and carbon monoxide with a comparable yield of products to other synthesis processes using a much larger concentration. Furthermore, the contact time between catalyst and reactants in the reaction zone is less than 5 seconds per pass, usually between about 0.5 and about 3 seconds. Contact time is defined as the time of contact of the synthesis gas with catalyst per pass. To effect the reaction in the presence of a finely-divided iron catalyst present in the reaction mixture in the amount heretofore described, according to an embodiment of this invention a reaction mixture of hydrogen and carbon monoxide is passed upwardly through an elongated, substantially vertical, reaction zone at a velocity greater than 8 feet per second and as high as 40 feet per second, although velocities as low as 5 or 6 feet per second may be used under certain conditions without departing from the scope of this invention.

At such velocities, the finely-divided hydrogenating catalyst, such as iron, is entrained or suspended in the reaction mixture in an amount between about 1 and about 18 pounds per cubic foot of gas and forms a continuous catalyst phase in the reaction zone. In some cases a concentration as high as 25 pounds per cubic foot is desirable but preferably lower concentrations are used. The reaction mixture and the catalyst which is entrained in the flowing gases are passed through the elongated reaction zone and are removed from the upper portion thereof together after less than about 5 seconds of residence therein. According to this invention, the catalyst is entrained in the reaction mixture in the reaction zone and flows therethrough in the direction of flow of the gases under conditions such that the conventional dense, pseudo-liquid catalyst phase is not formed.

In conventional fluid-type operations the catalyst forms a so-called "dense phase catalyst bed" in the reaction zone and consequently remains largely in the reaction zone itself until removed. Actually, two phases are formed in the reaction zone, a dense catalyst phase in the lower portion and a dilute phase having only a small amount of catalyst in the upper portion. The concentration of catalyst in such a dense phase is at least 25 pounds or 35 pounds per cubic foot of gas and usually between 50 pounds and 120 pounds per cubic foot.

It has been found in operating with a continuous catalyst phase according to this invention that a hydrogen to carbon monoxide feed ratio as low as 0.7 to 1 can be used with success. This ratio will vary from about 0.7 to 1 to about 1.5 to 1 and higher ratios may be used, if desired,

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but in any event the ratio is less than that generally required in conventional fluid-bed operations.

The catalyst employed in the present invention is a finely-divided powdered catalyst of a metal or metal oxide which is or becomes in the reaction zone a catalyst for the hydrogenating reaction. Finely-divided metallic iron or iron oxide or a mixture of metallic iron and iron oxide are an example of the catalyst employed in this invention. Preferably, a metallic iron catalyst is used in the finely-divided form. Other metals and metallic oxides may be employed which are effective in catalyzing the hydrogenation of carbon monoxide, such as cobalt, nickel, and other metals of group VIII of the periodic table. While the catalyst powder usually consists of such catalytic metals or their oxides, it may also include a minor amount of promoting ingredients, such as alkalis, alumina, silica, titania, thorium, manganese oxide, and magnesia. Also, the catalyst may be supported on a suitable support, such as a bentonite type clay, silica gel, "Super Filtrol" and mixtures of these supports. In the following description, catalyst powders consisting of a metal and/or a metal oxide and containing at most a minor proportion of promoters are referred to as finely-divided metal catalyst.

The exact chemical condition of the catalyst in its most active form is not certain. It may be that the active form is present when the metal is at an optimum degree of oxidation and/or carburization; consequently, a metallic iron catalyst which is in a reduced condition when first contacted with the reactants may reach its state of highest activity through being oxidized and/or carburized in the reaction zone. Therefore, in this specification and claims, the catalyst employed is described by reference to its chemical composition when first contacted with the reactants.

The catalyst is employed in a fine state of subdivision. Preferably, the powdered catalyst initially contains no more than a minor proportion by weight of material whose average particle diameter is greater than 250 microns. The greater proportion of the catalyst mass, preferably, comprises a material whose average particle diameter is smaller than 100 microns including at least 25 weight per cent of the material in a particle size smaller than 40 microns. An example of a desirable powdered catalyst is one which comprises at least 75 per cent by weight of material smaller than 150 microns and at least 25 per cent by weight of materials smaller than 40 microns.

The temperature of reaction for the hydrogenation of carbon monoxide is generally between about 300° F. and about 750° F. With a metallic iron catalyst, temperatures between 450° F. and 750° F. are usually employed. With a cobalt catalyst usually a temperature below 540° F. is sufficient for the hydrogenating reaction. Generally, the pressures employed are somewhat above atmospheric and range from about 10 pounds to as much as 500 pounds per square inch gage, preferably between about 80 pounds and about 300 pounds per square inch gage.

In effecting the reaction it may often become necessary to cool the reaction zone to maintain a relatively constant temperature. Various methods of cooling the reaction zone itself, such as by external cooling means or by injection of a cooling medium directly into the reaction mix-

ture, may be practiced without departing from the scope of this invention. Furthermore, it may often become necessary to preheat the reaction mixture prior to entry into the reaction zone, and also the catalyst may be preheated before introduction into the reaction mixture. However, the cooling and preheating are factors which will be characteristic of the particular apparatus being used and the particular conditions under which the reaction is effected.

Generally, the reaction zone itself will comprise a single or a multiple number of conduits or tubes of an inside diameter between about 1 inch and about 6 inches. Preferably the diameter of the reaction tubes is between about 1 inch and about 2.5 inches. It is known that the diameter of the tube is of considerable importance when entraining a catalyst in the reaction mixture, since the wall effect of the tube itself has a considerable effect on the disposition of the catalyst in the reaction stream. Consequently, it has been found that generally the reaction tube size should be below about 2.5 inches in diameter for optimum operations. However, with certain catalysts and with certain conditions of operation, tubes or reaction zones much larger in diameter may be used. From the standpoint of cooling the reaction zone, smaller tubes are also desirable since they present larger heat transfer surfaces.

According to a preferred embodiment of this invention, a fresh feed gas having a hydrogen to carbon monoxide ratio higher than the ratio in which these compounds are converted to other compounds is employed and the ratio of hydrogen to carbon monoxide in the reaction zone itself may be increased above the ratio in the feed gas and to a desired value by recycling a portion of the unconverted gas from the reaction zone, after removal of a part or all of the normally liquid product by condensation. A ratio of hydrogen to carbon monoxide in the fresh feed gas is used in which only a portion of the hydrogen is converted to products of the process. A portion of the effluent after removal of the greater part of the liquid product is recycled to the reaction zone in a volumetric ratio of recycle to fresh feed gas of about 0.5:1 to about 10:1, generally about 1:1 to about 5:1 or 6:1. The ratio of hydrogen to carbon monoxide in the reaction zone itself is usually about 1:1 to about 3:1 and according to this process may be maintained at about 1:1 without detrimental effect on the synthesis reaction. The ratio of hydrogen to carbon monoxide in the fresh feed itself may be considerably lower than in the reaction zone and usually ranges from about 0.7:1 to about 1.5:1. For very low feed ratios of less than 0.9:1 the ratio in the reaction zone itself may be even less than the feed ratio as a result of the consumption of hydrogen relative to carbon monoxide being greater than the feed ratio. Once through operations without recycle, although not generally used, are within the scope of this invention.

The linear velocity of the gaseous reaction mixture passing upwardly through the reaction zone is conveniently expressed in terms of superficial velocity, which is the linear velocity the feed stream would assume if passed through the reactor in the absence of catalyst, and takes into account the shrinkage in volume caused by the hydrogenation reaction. As previously stated, these superficial velocities are above about 8 feet per second, preferably above about 12 feet per second, and may be as high as 40 feet per second

or higher without departing from the scope of this invention. The superficial linear velocity is calculated from the arithmetic average of the gas rate at the bottom and top of the reaction zone.

The latter is arrived at by correcting the outlet gas volume for water and hydrocarbons condensed in the receivers, with corrections for pressure and average catalyst temperature. Contact times are calculated by dividing the length of the reaction zone by the superficial velocity.

The concentration of the catalyst in the gaseous reaction mixture in the reaction zone is usually less than about 18 pounds per cubic foot of gas, and preferably between about 3 pounds and about 12 pounds per cubic foot. The actual concentration required in the above range will depend to a certain extent upon the amount of inert gas in the reaction zone and also upon the accumulation of carbon and wax on the catalyst particles as the operation proceeds. The accumulation of wax and carbon on the catalyst decreases the weight of catalyst per cubic foot of gas thus the above values represent the extreme limits and may vary in accordance with this discussion.

Although the invention has been described with reference to an upwardly-flowing gaseous stream of reactants and catalyst, it should be understood that the catalyst and reactants may flow together downwardly, horizontally, or even angularly, through a reaction zone without departing from the scope of this invention. Important requirements of this invention are that the weight of the catalyst per cubic foot of gas and the contact time be within the limits previously discussed. It has been found that by upward flowing of gas through a substantially vertical reaction zone the weight of catalyst per cubic foot of gas and the reaction time can be controlled conveniently and accurately and for one reason is the preferable method of operation.

In operating a synthesis process according to this invention with an iron catalyst and at a temperature between about 550° F. and about 650° F. at relatively low super-atmospheric pressures a contraction of about 25 to about 85 per cent has been observed. The carbon monoxide disappearance is about 70 per cent to about 88 per cent and the selectivity of the reaction illustrated by the conversion of carbon monoxide to carbon dioxide is about 15 per cent to about 30 per cent. Condensed oil and water yields of about 30 to about 100 and about 80 to about 175 cc. per cubic meter of fresh feed gas, respectively, are obtained by operating according to the present process and may contain appreciable quantities of organic chemicals.

Certain particular advantages of the present process have been observed. One of these advantages which is of great importance is the fact that a relatively low hydrogen to carbon monoxide feed ratio can be utilized and as a result a proportionate decrease in the cost of the production of hydrogen is effected. At the extremely high velocity capable of being used by the present process and apparatus much less catalyst is used and much greater efficiency of heat transfer is obtained. Furthermore, temperature conditions can be easily and readily controlled at such high velocities and at such short contact times. By the use of a short contact time higher selectivity can be obtained as the result of decreased side reactions, such as over-polymerization. Less than 3 per cent per week wax accumulation on the catalyst has been observed with

the present method which compares with as much as 23 per cent per week wax accumulation with a low velocity fluid-bed type operation. Carbon deposits upon the catalyst have also been observed to be much less than those observed in low velocity operations. A carbon deposit of less than about 2 per cent per week for the present process as compared to above about 5 per cent per week for the low velocity process has also been observed.

As a result of high velocity and increased heat transfer, it is possible to use much greater temperature gradients between the catalyst and a cooling medium, if one is used, with the present process. In the low velocity fluid-bed operation a temperature difference between coolant and reaction mixture between 10° F. and 100° F. is conventional. However, with the present process a gradient considerably in excess of 100° F. is possible. Also, relatively cold gas feed may be employed to the reaction zone. Mixing is so rapid under such conditions that no deleterious effect is observed.

The invention will be described further by reference to the accompanying drawings which are views in elevation, partly in cross-section, of suitable apparatus for carrying out the process of the present invention. Fig. 1 of the drawings is an elevational view diagrammatically illustrating a reaction zone and suitable auxiliary equipment for carrying out one embodiment of the present invention. Figures 2 and 3 of the drawings are other reaction chambers embodying the essential features of the present invention and may be substituted for the reaction chamber shown in Fig. 1 of the drawings.

In Fig. 1 of the drawings a synthesis gas comprising hydrogen and carbon monoxide present in a ratio between about 0.7:1 and about 1.4:1 is obtained from any suitable source. For example, a suitable source of hydrogen and carbon monoxide is the conversion of steam, carbon dioxide, and methane in the presence of a suitable catalyst, such as nickel. The resulting mixture of such a conversion usually contains sulphur and sulphur compounds, and the gas is preferably purified to remove such compounds therefrom. If a sulphur resistant catalyst is used the purification step is unnecessary. After purification in conventional manner known to those skilled in the art, the mixture of hydrogen and carbon monoxide is introduced into the lower end of a 26 foot length 1 inch diameter conduit or tubing 8 of Fig. 1. Conduit 8 is a curved conduit which has a major portion thereof positioned substantially vertically. Conduit 8 is also lagged. The gaseous reaction mixture is passed upwardly through conduit 8 and catalyst from a standpipe 10 is introduced into the flowing gaseous stream in the lower portion of conduit 8, as shown. The velocity of the gas in conduit 8 is maintained above about 8 feet per second in the vertical section in order to prevent the formation of a pseudo-liquid dense phase of catalyst in the vertical section of the conduit, but instead to form a continuous catalyst phase of relatively dilute concentration. Usually the velocity of the gaseous stream in conduit 8 is between about 12 and about 40 feet per second and at such velocities the catalyst is entrained in the gaseous stream and passes overhead into the upper portion of standpipe 10 with the gaseous stream itself. Since a continuous catalyst phase is present in conduit 8, the amount of catalyst taken overhead into the standpipe is approxi-

mately equivalent to the amount of catalyst introduced into the lower portion of conduit 8 from standpipe 10.

Standpipe 10 comprises two substantially vertical, concentric, extra heavy steel pipes; an outside pipe 11 and an inside tubing 10. The outside pipe 11 is welded at its ends to the inside tubing 10 to form an enclosing jacket which may be filled with a liquid as a cooling or heating medium. The cooling or heating medium is introduced into the annular space formed by the two concentric pipes through line 12 and may be withdrawn therefrom through line 15. In some instances the cooling or heating medium may be introduced through line 15 and removed through line 12, if desired. In another embodiment where the liquid introduced into the annular space is evaporated therein and the latent heat of evaporation is used to cool the catalyst, liquid is introduced through line 12 and vapors are removed, also through line 12.

Catalyst passes from conduit 8 into a conical section 14 which has a larger diameter than conduit 8 and thereby the velocity of the gases are diminished and the catalyst separates from the gaseous stream and flows downwardly into conduit 10. A slide valve 13 in the lower portion of standpipe 10 regulates the flow of catalyst from conduit 10 into conduit 8. The upper end of conduit 10 is connected by means of a conical section 14 to an enlarged conduit 16. Conduit 16 facilitates the disengagement of the catalyst from the gas stream after the passage of the latter into conical section 14. Conduit 16 is connected by means of manifold 17 with conduits 18 and 19. Conduits 18 and 19 contain ceramic filters 20 and 21 which are constructed of porous material, such as alundum, permeable to the gases and vapors emerged from conduit 8 but impermeable to catalyst "fines" entrained in the gaseous effluent. Filters 20 and 21 are cylindrical and are closed at the bottom ends. A substantial annular space is provided between the wall of the filters and the wall of the enclosing conduit for the passage of gases and vapors and entrained catalyst upwardly through the annular space between the filters and conduits 18 and 19. The upper ends of filters 20 and 21 are mounted inside conduits 18 and 19 by means of enclosure means 22 and 23. The gases and vapors must pass through either or both filters 20 and filter 21 to reach outlet conduits 24 and 26.

Union of the various conduits in the upper portion of the standpipe 10 is made by welding.

When using a metallic iron catalyst in a finely-divided state, usually between about 40 and about 150 microns, the temperature of reaction in conduit 8 is between 550° F. and about 650° F. A pressure of about 80 pounds per square inch gage has been found to be quite satisfactory. However, various pressures above and below this may be used without departing from the scope of this invention. With a velocity greater than 10 feet per second in conduit 8 the reaction time is less than 3 seconds per pass. Operating at a gas velocity of about 15 feet per second in conduit 8 a loading of about 6 pounds of catalyst per cubic foot of gas entering conduit 8 will produce a concentration of about 12 pounds of catalyst per cubic foot of gas in the vertical section of conduit 8. Similarly, if the loading is cut to about 4 pounds per cubic foot of gas the concentration of catalyst in the vertical section of conduit 8 is about 8 pounds per cubic foot. In all cases the velocity of the gas passing

through conduit 8 is maintained above about 5 or 6 feet per second in order to prevent a formation of a dense pseudo-liquid phase of catalyst and to assure a continuous phase of catalyst in conduit 8. In operating at such high velocities the catalyst is entrained in the gaseous mixture and flows from the lower portion of conduit 8 to the upper portion thereof and settles or separates from the gaseous mixture in conical section 14 and enlarged conduit 16. The gaseous portion of the mixture from conduit 8 flows upwardly through the filters 20 and 21 and into the respective outlet conduits 24 and 26. Separated catalyst flows downwardly through conduit 10 and by the regulation of valve 13 is introduced into conduit 8 at the desired rate. The density of the catalyst in conduit 10 is usually about 50 pounds to 110 pounds per cubic foot of volume.

In some instances, the heat of reaction may be removed by cooling the catalyst and using the sensible heat of the catalyst as a means for cooling the reaction mixture in the reaction zone. To accomplish this end, the catalyst in this particular apparatus may be cooled in conduit 10 by introducing a liquid, such as water or Dowtherm, through conduit 12 into the annular space between concentric conduits 10 and 11. The evaporation of the water or Dowtherm in the annular space removes a large portion of the heat in the catalyst. The cooled catalyst is then introduced into conduit 8 for recycling through the reaction zone or conduit 8. Conduit 8 may be cooled directly itself by indirect heat exchange (not shown) without departing from the scope of this invention. Various other methods known to those skilled in the art may be used to cool either the reaction mixture in conduit 8 or the catalyst in conduit 10 without departing from the scope of this invention.

Since the pressure differential between just below slide valve 13 and the upper portion of reaction zone 8 may vary to a considerable extent, it is necessary to control slide valve 13 to compensate for the pressure differential in order to obtain a constant flow of catalyst from standpipe 10 into conduit 8. This control of slide valve 13 is obtained by connecting a differential pressure recorder 28 by means of conduits 27 and 29 to the upper and lower portions of standpipe 10, as shown, and transmitting changes in pressure differential to slide valve 13 by means of element 31 so that when the pressure differential is increased valve 13 is closed slightly and when the pressure differential is decreased valve 13 is opened slightly. The concentration of the catalyst per cubic foot of gas in the vertical section of conduit 8 may be determined by connecting a pressure differential recorder (not shown) on the vertical section of conduit 8 and calibrating the recorder readings in terms of concentration of catalyst. The gaseous effluent from either conduit 18 or 19 is passed through filters 20 and 21 into outlet conduit 24 and 26, respectively. Usually only one outlet conduit is used at a time. Thus, for example, the gaseous effluent passes through outlet conduit 24 to conduit 33, through condenser 34 where the effluent is cooled to about 40° F. at operating pressure and then passed to accumulator 36. In accumulator 36, gaseous components are separated from liquid components of the cooled effluent. Uncondensed components of the effluent, such as hydrogen, carbon monoxide, methane, propylene, butylene, light naphtha, and organic oxygenated compounds, are recycled by means of conduits 38 and 44 and a

compressor or blower 45 to the lower portion of conduit 8 in a ratio of about 1:1 to about 5 or 6:1 of volumes of recycle to volumes of fresh synthesis gas. The amount of unreacted hydrogen and carbon monoxide in the recycle gas determines how much the ratio of carbon monoxide and hydrogen in the reaction zone itself will deviate from the ratio in the original feed. As shown, the recycle gas is introduced into the fresh feed before the catalyst is introduced into the gaseous mixture, however, the recycle may be introduced into the gaseous mixture after the catalyst is introduced into the feed stream or the feed gas may be introduced into the gaseous mixture after the introduction of the catalyst without departing from the scope of this invention.

After passage of the gaseous effluent through filter 20 for a time, the filter becomes coated and clogged with catalyst fines which have not settled out from the gaseous effluent. In order to remove these fines from the catalyst filters so as to ensure continuous passage of the gaseous effluent through the filters and so as to recover the catalyst, the course of gaseous effluent is changed to flow through filter 21 and conduit 26 and a portion of the uncondensed effluent is passed from accumulator 36 by means of conduit 38, compressor 41, and conduit 24 to filter 20. The pressure of the gas blows the fines from the filter into conduit 18. The fines then settle in conduit 18 to standpipe 10. Other gases than the uncondensed effluent may be used to remove the fines from the filter and may be introduced through line 42, if desired.

Liquid condensate in accumulator 36 is passed through conduit 39 to separating means 48, which may represent various separating units, such as distillation columns, absorption units, extraction units, and the like. In separating means 48, water is separated from organic compounds and removed through line 49; oxygenated organic compounds are separated therein and removed through line 51; and hydrocarbons are separated and removed through line 52.

Uncondensed gas from accumulator 36 not used for recycle, etc., is removed from the system through conduit 40 and passed to oil and chemical recovery equipment not shown.

It has been found that operating a synthesis process according to this invention, in which the synthesis gas is passed through a reaction zone at a high velocity, good yields of products are realized. Ordinarily one would believe that insufficient catalyst for accomplishing the desired reaction would be carried by the gas at such high velocities, but it has been found that within the range indicated sufficient catalyst is carried by the gas to effect the reaction between hydrogen and carbon monoxide. If desired, the synthesis gas entering conduit 8 may be preheated; but it has been found that preheating the gas is unnecessary in most instances and that the contact of the hot catalyst from standpipe 10 with fresh feed gases entering conduit 8 does not cause balling or agglomeration of the catalyst mass. It has also been found, as previously mentioned, that the wax and carbon content of the catalyst with extended use is much less than that observed in the conventional fluid-bed operations, and it is believed the reason for this is that catalyst eddy-currents are minimized in high velocity operations which greatly shortens the contact time between catalyst and reactants per pass. Longer catalyst life may also result from shorter

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contact time per se between reactants and catalysts. The above theory is offered merely as a possible explanation of the extended life of the catalyst realized in the present invention and is not considered to unnecessarily limit the invention.

Fig. 2 of the drawings is another arrangement of apparatus suitable for carrying out this invention. The apparatus shown in Fig. 2 may be substituted for conduit 8 and standpipe 10 of Fig. 1. The filter sections of Figures 1 and 2 are the same. Accordingly, a synthesis gas comprising hydrogen and carbon monoxide is passed into a reaction zone 70 through a line 68. Reaction zone 70 comprises a bundle of substantially vertical tubes 71 through which the gases pass upwardly into a cylindrical section 74 of a larger total cross-sectional area than the total cross-sectional area of tubes 71. The outer surface of tubes 71 are sealed off to form an annular space 72 between the inner surface of the outer shell of reaction zone 70 and the outer surface of tubes 71. A cooling (or heating) fluid may be passed through annular space 72 to cool (or heat) the reaction mixture as it passes upwardly through tubes 71. Tubes 71 correspond to conduit 8 of Fig. 1. Upon reaching enlarged section 74 the velocity of the gaseous mixture is decreased to such an extent that the catalyst settles from the effluent and passes down through a standpipe 73 into the lower portion of reaction zone 70 where the catalyst falls or is drawn into a high velocity gaseous stream passing upwardly into tubes 71. The lower portion of reaction zone 70 and tubes 71 are of such a cross-sectional area that the catalyst is entrained in the gaseous stream.

The cooling medium is introduced into the annular space 72 through line 76 and is withdrawn therefrom through line 77.

The reaction effluent from reactor 70 passes upwardly through conduits 17, 18, and 19 of which the latter two contain filters as previously discussed with reference to Fig. 1. Conduits 17, 18, and 19 are the same as conduits 17, 18, and 19 of Fig. 1. The gaseous effluent passes from conduits 18 and 19 through outlet conduits 24 and 26 to conduit 33. The effluent in conduit 33 of Fig. 2 is condensed and separated according to the description of Fig. 1 and a portion of the uncondensed gases may be recycled (not shown) to conduit 68, if desired.

The cross-sectional area of conduits 71 with respect to the quantity of gases flowing there-through is such that the velocity of the gases is greater than about 8 feet per second, while the cross-sectional area of enlarged section 74 is such that the velocity of the gases is below about 5 feet per second so that the catalyst may settle from the gaseous effluent. In section 74, the catalyst may form a dense pseudo-liquid catalyst phase having a concentration of catalyst greater than about 20 pounds or 25 pounds per cubic foot of gas. In such a case, reactor 70 has two reaction sections, tubes 71 in which the gas flows at a relatively high velocity and with a relatively low concentration of catalyst and enlarged section 74 in which the gas flows at a relatively low velocity and with a relatively high concentration of catalyst. The same effect may be achieved in the apparatus of Fig. 1 of the drawings by extending the length of section 16 and adjusting the cross section thereof such that the velocity of the gases therein are appropriate for the formation of a dense pseudo-liquid catalyst phase. In

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this manner, a high velocity continuous catalyst phase will exist in conduit 8 and a dense pseudo-liquid catalyst phase will exist in section 16.

A moveable valve 69 is provided at the lower end of standpipe 73 to control the flow and dispersion of the catalyst into the gaseous stream. Valve 69 may cause an aspiration effect by the deflection of the gases passing by and as a result of which the catalyst is drawn into the gaseous stream.

Fig. 3 gives a diagrammatic illustration in elevation of another arrangement of apparatus for the synthesis of hydrocarbons according to the present invention. The apparatus in Fig. 3 is very similar in operation to the apparatus of Fig. 1 and Fig. 2 and thus only a brief discussion of its operation will be included. A synthesis gas passes through a conduit 104 to a reaction chamber 106. Reaction chamber 106 comprises a bundle of reaction tubes 107 surrounded by a shell 108 to form an annular space 109 between the outside diameter of tubes 107 and shell 108. Annular space 109 is for the circulation of cooling fluid around reaction tubes 107 in order to maintain the temperature of reaction substantially constant. The cooling fluid, such as Dowtherm, enters annular space 109 through conduit 113. The pressure in annular space 109 is such that the Dowtherm boils below the desired temperature of reaction in tubes 107. The vaporized Dowtherm passes from annular space 109 through conduit 112 to an accumulator 114. From accumulator 114 vapors pass through line 115 to a condenser 116 in which substantially all of the Dowtherm vapor is condensed. Condensate from condenser 116 passes to accumulator 114 through conduit 118. From accumulator 114 liquid Dowtherm is recycled to annular space 109 through conduit 113. Any uncondensable vapors are removed from the system through conduit 117.

A reaction effluent comprising reaction products and finely-divided catalyst entrained in the reaction effluent is passed to a settling and accumulation chamber 120 through conduit 111. Chamber 120 comprises an upper settling chamber 121 and a lower accumulation chamber 128. The cross-sectional area of settling chamber 121 is such that substantially all of the catalyst separates from the gaseous effluent and flows downwardly through a funnel-shaped septum 127 into the lower portion of accumulation chamber 128. Gases containing a small amount of fine catalyst pass upwardly in settling chamber 121 into a cyclone separator 122. Gases from cyclone separator 122 are removed therefrom by conduit 123 and may be treated in the manner heretofore described. Separated fine catalyst is removed from cyclone separator 122 by means of conduit 124 and passed to funnel-shaped septum 127, as shown. Finely-divided catalyst accumulates in accumulation chamber 128 to a level above the end of the funnel 127 in order to prevent the passage of the gaseous effluent downwardly through funnel 127. An aeration gas, such as hydrogen or recycle gas, is introduced into the lower portion of accumulation chamber 128 through conduit 132 and is injected into the accumulated catalyst therein by means of dispersion means 133 which may comprise a perforated conduit or the like. Accumulator 128 may be maintained at a substantially higher pressure than settling chamber 121 by introducing a gas therein through conduit 131, if desired. By maintaining the pressure higher in chamber 128 than in chamber 121 passage of gaseous effluent

into the accumulated catalyst is prevented. Pressuring gas and aeration gas may be passed, if desired, from chamber 128 to chamber 121 through conduit 134. Alternatively, aeration gas may be passed from chamber 128 through conduit 131 if that conduit is not being used for the introduction of a pressuring gas.

Accumulated catalyst in accumulator 128 passes through a standpipe 129 to conduit 104 to be mixed with fresh feed.

Cyclone separator 122 may be omitted and in its place another type of separating means may be used, such as the filter means of Figures 1 and 2. Various other modifications of Fig. 3 may become obvious to those skilled in the art without departing from the scope of this invention.

Assuming a reactor inlet gas volume of 25,000 standard cubic feet per hour, a reactor consisting of four tubes having an inside diameter of 2 inches would operate at a maximum linear velocity of 9 feet per second. Using a tube length of 36 feet, a velocity of 9 feet per second would correspond to about 4 seconds of contact time per pass, allowing for contraction. A reactor capable of a 40 feet per second linear velocity and 5 seconds contact time per pass would involve a flow path of about 200 feet in length. A single tube having a 2 inch inside diameter might be substituted for the four tubes having a 1 inch inside diameter with substantially the same velocity and throughput.

Various modifications and alterations of the apparatus and flow shown in Figures 1, 2, and 3 may be practiced by those skilled in the art without departing from the scope of this invention. Various coolers, condensers, distillation units, and other separating means have not been shown for a matter of convenience and simplicity, but their presence will be obvious to those skilled in the art.

The following example is offered as a means of better understanding the application of the present invention to the hydrogenation of carbon monoxide and the specific recitation of certain limitations therein is not considered unnecessarily limiting to the present invention.

EXAMPLE

In this example an iron catalyst was employed, which iron catalyst was prepared by fusing and subsequently reducing Alan Wood ore. The catalyst contains metallic iron, alumina, titanium dioxide, and silica, and about 1.2 to about 1.4 weight per cent potassium calculated as the oxide. The potassium oxide is incorporated with the catalyst prior to fusion. The particle size of the catalyst employed in the runs of this example is shown in Table I below:

Table I

POWDERED IRON CATALYST PARTICLE SIZE

Roller Analysis	Weight Percent
Microns:	
0-10.....	16.2
10-20.....	17.0
20-40.....	19.2
40-60.....	24.0
60+.....	23.6
Recovery, percent.....	98.8
Density (basis water).....	6.8

Having prepared a catalyst of the desired properties and the required size, the catalyst was

introduced into a conduit, similar to conduit 8 of Fig. 1 of the drawings, in which a mixture of hydrogen and carbon monoxide was flowing upwardly at a relatively high velocity. The conditions of reaction and the analysis of the product is shown in Table II below. In substantially all respects the apparatus used for obtaining the data shown in Table II was the same as the apparatus of Fig. 1 of the drawings. In obtaining data various gas velocities as well as concentrations of catalyst in the reaction zone (conduit 8) were used to determine their effect on the reaction and product. Also it will be noted that the pressures used varied from 80 to 150 pounds per square inch gage. In some runs, especially the later runs, the temperature of the inlet feed gas was decreased to a relatively low value to determine the effect of contact between relatively hot catalyst and relatively cold gas and whether temperature conditions could be maintained in the reaction zone. The reaction effluent was withdrawn from the reaction zone after a relatively short contact time with the catalyst. The catalyst separated by gravity from the effluent in an enlarged section of the apparatus and passed by means of a standpipe, such as conduit 10 of Fig. 1, to the point of introduction of the catalyst into the synthesis gas stream. The effluent passed through filters to separate fine catalyst therefrom. The filters were cleaned intermittently or continuously by flowing recycle gas back through them as previously explained with reference to Fig. 1 of the drawing. The effluent then passed through a condenser at 40° F. and at operating pressure and uncondensed gases were recycled to a point just before the first contact of catalyst and synthesis gas.

Catalyst in the standpipe was maintained at about 600° F. by means of electrical heating elements wound around the 4-inch jacket surrounding the 2 inch standpipe. Various predetermined loadings of the catalyst into the synthesis gas stream were used and the tendency for variation in loadings for any particular run caused by variation in differential pressure in the reaction zone was minimized by controlling a slide valve on the bottom of the standpipe by a conventional differential pressure recorder responsive to the differential pressure between the top and bottom of the standpipe. The differential pressure recorder was set at various readings of inches of water depending upon the catalyst loading desired. The concentration of catalyst in the reaction zone or elongated conduit was determined by the differential pressure in a 15 foot vertical section of the conduit. The overall length of the reaction zone was about 26 feet and 3½ inches, i. e., from the point the catalyst was first contacted with the synthesis gas and the point where the catalyst was separated from the reaction effluent.

The velocity of the gas stream was so great that the catalyst was entrained in the synthesis gas stream throughout the reaction zone in a continuous phase without the formation of the conventional dense pseudo-liquid phase of catalyst.

Gas bleeds into the instrument lines were used to prevent catalyst from clogging instrument lines and instruments. In some cases recycle gas was used as the bleed gas, in other instances hydrogen or fresh feed gas were used.

The catalyst in the standpipe was aerated with recycle gas or combined inlet gas in most instances, however other gases, such as carbon dioxide, hydrogen, nitrogen, and steam could have been used if desired.

Table II

Run No.	1	3	4	5	6	7	9	12	13	14	15	18	19	20	21	24	25
Hours on run	6	6	16	18	18	150	12	24	22	6	18	18	24	24	24	48	24
Operating conditions:																	
Pressure, p. s. i.	80	80	80	80	80	150	150	150	150	150	150	150	150	150	150	150	150
Ratio of recycle to fresh feed	6.0	7.7	1.7	1.8	6.2	4.92	4.7	8.5	6.5	8.1	6.6	6.5	6.4	5.6	5.4	6.6	5.8
Fresh feed, C. F./H.	88	143	150	155	191	251	224	106	133	105	127	135	128	145	148	126	171
Gas ratio, H ₂ :CO—																	
Fresh feed	3	3	3	3	3+	3	2	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4
Inlet to reaction zone				4.5	14	4.9	3.9	2.0	2.2	2.0	2.2	2.6	2.6	1.7	2.0	2.1	—
Standpipe:																	
Temp., ° F.	593	598	598	596	605	597	597	596	597	601	600	604	598	600	595	559	591
Density, P. C. F.	97	100	103	99	102	106	87	87	82	80	78	69	67	67	66	55	49
Lin. velocity, F. P. S.	0.4	0.4	0.10	0.17	0.18	0.14	0.13	0.11	0.11	0.08	0.10	0.10	0.09	0.09	0.11	0.10	0.12
Reaction zone:																	
Temp., ° F., in zone	585	595	594	596	603	601	600	600	604	604	602	602	595	590	585	540	602
Gas inlet temp., ° F.	572	616	604	603	605	576	603	600	600	600	600	550	500	450	400	150	615
Line velocity, F. P. S. (superficial)	15	19	7	8	15	14.3	13.4	10.5	10.4	10.2	10.3	10.6	9.8	8.9	9.5	8.5	10.1
Catalyst loading (inches of water)	35	55	53	35	49	70	70	70	70	75	80	59	58	59	58	57	—
Concentration of catalyst, P. C. F.	3.5	5.9	3.5	3.0	4.9	8.4	8.0	8.7	9.4	10.1	11.2	8.2	8.4	8.7	8.0	8.0	5.8
Contact time, sec.	1.7	1.4	3.7	3.2+	1.7	1.8	1.9	2.5	2.5	2.5+	2.5	2.7	2.7	2.9	2.7+	3.1	2.6
Results:																	
Contraction, percent	—	54	26	23	82	35	37	35	46	37	43	46	49	49	44	43	62
CO disappearance	85	79	67	54	81	88	67	71	74	83	83	—	—	79	—	—	—
Observed oil, cc./m. ³	20	19	28	31	44	38	41	101	80	93	80	63	70	69	—	50	98
Water, cc./m. ³	167	105	81	80	—	135	110	165	142	168	129	141	160	121	—	95	154
Selectivity—																	
CO→CO ₂ , percent	12.5	14.7	22.4	15.4	—	13.2	20.5	23.2	23.2	—	11	17.4	17.4	20.0	—	17.9	—
CO→CH ₄	—	—	—	—	—	—	—	18	14.5	—	17.8	12.9	12.9	7.4	—	—	—

From the above data it is apparent that the product and results in general compare favorably with fluid-bed operations using a conventional dense phase of catalyst. It should be noted that a considerably lower ratio of hydrogen to carbon monoxide in the feed gas is used than is usually practiced in fluid-bed operations. Since the concentration of catalyst is so small as compared to other type operations, the amount of catalyst per unit volume of gas throughput is correspondingly small.

It is assumed that a catalyst slip of about 50 per cent is present in the vertical section of the reaction zone. Actually, therefore, the catalyst loading of pounds of catalyst per cubic foot of gas may be about half the concentration of catalyst in the reaction zone. Thus, for a concentration of 8 pounds per cubic foot of gas, the catalyst flow was calculated to be about 1000 pounds per hour.

When operating at 1.4 ratio of hydrogen to carbon monoxide, as in runs 13 to 15 and 16 to 21, with a recycle ratio of about 5:1 to 6:1 the gas inlet composition (including fresh feed and recycle gas) was about 18 per cent carbon monoxide and a ratio of H₂:CO of about 2:1 to 2.5:1 existed in the reaction zone.

In certain runs, it was found that with a catalyst loading represented by a differential pressure of 80 inches of water the flow of catalyst in the reaction zone was too great to permit uniform settling in the standpipe. However, with a larger diameter or longer standpipe or bigger slide valve, loadings higher than that represented by 80 inches of water could be used satisfactorily. In run 24 a combined inlet gas temperature of 150° F. was used. When operating at a combined recycle and fresh feed gas temperature as low as 150° F. it was necessary to maintain the catalyst temperature at least 560° F. to ensure a sufficiently high reaction temperature. The catalyst was maintained at about 560° F. in the standpipe by heating the catalyst electrically, as previously discussed. With the present equipment, the catalyst could not be maintained at a high enough temperature to permit a lower inlet gas temperature. It is believed that an inlet gas temperature as low as

100° F. or lower could be used if the catalyst could be maintained at 600° F. in the standpipe.

In runs Nos. 20 and 21 the selectivity appeared to be good with 21 per cent CO→CO₂ and a small amount of CO→CH₄. Analysis of the product of run 20 indicated that about 40 per cent of the condensed oil comprised oxygenated compounds of which 15 per cent were acids. The water contained about 14 per cent chemicals, approximately half of which were acids. A yield of about 40 cc./m.³ of total oxygenated compounds was obtained.

Operations at 250 pounds per square inch pressure have proved to be successful with results somewhat similar to those at lower pressures.

As will be noted, no external cooling of the reaction system was used during the present set of runs for the hydrogenation of carbon monoxide by the process of this invention. In fact, heat was added to the system through the catalyst standpipe by electrical heating coils around the jacket of the standpipe which contained Dowtherm under pressure. The reaction zone was heavily lagged and electrically wound to produce substantially adiabatic conditions in that zone. If the entire assembly of apparatus had been completely adiabatic, and if the reaction had been carried out on a larger scale, cooling of the standpipe or the reaction zone would probably have been necessary. Such cooling could have been carried out by contact with a cooling medium surrounding the reaction zone, such as cooling with water or Dowtherm, or by introducing relatively cold feed gas or cold recycle gas or both.

Prior to run 12 an iron catalyst similar to that previously described, but which had been used in conventional dense-phase, fluid-bed operation for about 400 hours, was substituted for the original catalyst in carrying out the process. The composition of this catalyst prior to use in the process is shown in column 1 of Table III. After the catalyst had been used for a period of two days, five days, and nine days, analysis of the catalyst was made to determine the effect of this type of reaction on the carbon and oil con-

tent of the catalyst, and also upon the catalyst size. This data is shown in Table III.

Table III

CATALYST ANALYSIS

	Charged	2 days	5 days	9 days
Chemical Analysis:				
Oil+Wax.....	4.0	4.5	5.5	5.2
Carbon.....	23.1	23.5	25.4	24.0
Total Iron.....	65.3	65.2	63.4	62.0
Roller Analysis:	Weight percent			
Microns—				
0-10.....		0.9	1.0	1.2
10-20.....		0.9	2.8	2.7
20-40.....		7.5	10.2	12.9
40-80.....		32.4	33.3	38.4
80+.....		58.3	52.7	44.8
Density.....		3.4	3.1	3.2

The data of Table III indicates only a slight change in chemical composition and size of the catalyst from the original material introduced at the beginning of run 12. The wax content increased about 1.5 per cent and the fixed carbon content increased about 2.3 per cent. A decrease of less than 2 per cent in the total iron was noted. The particle size analysis indicates that the catalyst became slightly finer on use. From these results the catalyst composition in the circulating system of the present invention was probably more stable than with the conventional dense-phase operation using a fluid-bed.

This invention has been described with particular reference to specific apparatus and to specific conditions of operation. However, various modifications of the apparatus and various operating conditions may be used within the limits disclosed without departing from the scope of this invention. Essentially, the apparatus design itself must permit high velocity of gases in at least a portion of the reaction section.

Having described our invention, we claim:

1. A process for the hydrogenation of an oxide of carbon to produce organic compounds having more than one carbon atom per molecule which comprises introducing a finely-divided metal hydrogenating catalyst comprising iron and a gaseous mixture comprising hydrogen and an oxide of carbon into the lower portion of an elongated reaction zone, passing gases upward through said elongated reaction zone at a linear velocity of at least about 5 feet per second to suspend said catalyst in the gases such that the concentration of catalyst in gases is smaller than the concentration of the same catalyst when utilized under operating conditions to obtain a dense phase catalyst bed and such that finely-divided metal catalyst moves in the direction of flow of the gases through said elongated reaction zone, maintaining suitable conditions of temperature, pressure and residence time for the conversion of a major proportion of said oxide of carbon in said elongated reaction zone, maintaining the temperature of said reaction zone at at least 550° F. and within a relatively narrow range of not greater than 100° F. by indirect heat exchange with a cooling medium which is at a temperature substantially lower than the temperature of reaction, passing from the upper portion of said elongated reaction zone a gaseous effluent containing finely-divided metal hydrogenating catalyst through a confined passageway to a separation zone in which finely-divided catalyst is separated from gases, maintaining the pressure drop in the confined passage-

way between said reaction zone and said separation zone not greater than about 80 inches of water, recovering from said effluent organic compounds having more than one carbon atom per molecule as products of the process, withdrawing catalyst from said separation zone, and passing catalyst thus separated to the lower portion of said elongated reaction zone at a temperature within about 20° F. of the reaction temperature through a confined passageway.

2. The process of claim 1 in which the temperature of the cooling medium is at least 100° F. below the temperature of reaction.

3. A process for the hydrogenation of carbon monoxide to produce organic compounds having more than one carbon atom per molecule which comprises introducing a finely-divided hydrogenating catalyst comprising iron and an alkali as a promoter and a gaseous mixture comprising hydrogen and carbon monoxide into the lower portion of an elongated reaction zone, passing gases upward through said reaction zone at a linear velocity of at least about 5 feet per second to suspend said catalyst in the gases such that the concentration of catalyst in said gases is smaller than the concentration of the same catalyst when utilized under operating conditions to obtain a dense phase catalyst bed and such that finely-divided catalyst moves in the direction of flow of said gases in said elongated reaction zone, maintaining a temperature of reaction between about 550 and about 650° F. and a pressure between about 10 and about 500 pounds per square inch gage such that a major proportion of the carbon monoxide is converted, controlling the temperature of reaction by indirect heat exchange with a cooling medium which is at a substantially lower temperature than said reaction temperature, removing from the upper portion of said elongated reaction zone a gaseous effluent containing finely-divided catalyst and passing same through an unrestricted confined passageway whereby the pressure drop in said passageway is not greater than about 80 inches of water to a separation zone in which catalyst is separated from gases, recovering organic compounds having more than one carbon atom per molecule from the gaseous effluent as products of the process, withdrawing catalyst from said separation zone, and passing catalyst thus separated directly to the lower portion of said elongated reaction zone at a temperature not lower than about 560° F. through a confined passageway.

4. The process of claim 3 in which the linear gas velocity is between about 8 and about 40 feet per second.

5. A process for the hydrogenation of carbon monoxide to produce organic compounds having more than one carbon atom per molecule which comprises introducing a finely-divided metal hydrogenating catalyst comprising iron and a gaseous mixture comprising hydrogen and carbon monoxide into the lower portion of an elongated reaction zone, passing gases upward through said elongated reaction zone at a linear velocity of at least about 6 feet per second to suspend said catalyst in the gases and such that finely-divided catalyst continuously moves in the direction of flow of gases in said elongated reaction zone, maintaining a reaction temperature between about 550 and about 650° F. and a reaction pressure between about 10 and about 500 pounds per square inch gage such that a major proportion of the carbon monoxide is converted, maintaining the reaction temperature within a relatively

narrow range within the above temperature range by indirect heat exchange with a cooling medium which is at a temperature substantially below the temperature of reaction, removing from the upper portion of said elongated reaction zone an effluent containing finely-divided catalyst and passing same through a confined passageway to a separation zone in which catalyst is separated from gases, maintaining the pressure drop in said confined passageway between said reaction zone and said separation zone not greater than 80 inches of water, cooling and condensing said effluent to form a vapor phase and a liquid phase comprising organic compounds, recovering from said liquid phase organic compounds as products of the process, recycling at least a portion of the vapor phase to the lower portion of said elongated reaction zone in a volumetric ratio of recycle gas to fresh feed gas of about 0.5:1 to about 10:1, withdrawing catalyst from said separation zone, contacting catalyst thus separated with a hydrogen-containing gas, returning catalyst thus separated and contacted with a hydrogen-containing gas to the lower portion of said elongated reaction zone at a temperature not lower than about 560° F., controlling the rate of return of said separated catalyst to the lower portion of said elongated reaction zone such that catalyst loading rate is at least 4 pounds per cubic foot of gas and such that the concentration of catalyst into said elongated reaction zone is less than about 25 pounds per cubic foot.

6. The process of claim 5 in which said hydrogenating catalyst comprising iron contains an alkali and the pressure is between about 80 and about 300 pounds per square inch gage.

7. The process of claim 5 in which said catalyst concentration is between about 1 and about 18 pounds per cubic foot.

8. A process for the hydrogenation of carbon monoxide to produce organic compounds having more than one carbon atom per molecule which comprises introducing a finely divided hydrogenating catalyst comprising iron and an alkali as its promoter containing no more than a minor proportion by weight of material whose average particle diameter is greater than 250 microns and a gaseous mixture comprising hydrogen and carbon monoxide into the lower portion of an elongated reaction zone, passing gases upwardly through said elongated reaction zone at a linear velocity of at least about 5 feet per second to suspend said finely divided catalyst in the gases such that the concentration of catalyst in said gases is not greater than about 25 pounds per cubic foot of gas and such that finely divided catalyst moves in the direction of flow of said

gases in said elongated reaction zone, maintaining a temperature of reaction between about 550° F. and about 650° F. and a pressure between about 10 and about 500 pounds per square inch gage such that a major proportion of the carbon monoxide is converted, controlling the temperature of reaction by indirect heat exchange with a cooling medium which is at a substantially lower temperature than the reaction temperature, removing from the upper portion of said elongated reaction zone a gaseous effluent containing finely divided catalyst and passing same through an unrestricted confined passageway whereby the pressure drop in said passageway is not greater than about 80 inches of water to a separation zone in which catalyst is separated from gases, recovering organic compounds having more than one carbon atom per molecule from the gaseous effluent as products of the process, withdrawing catalyst from said separation zone, stripping catalyst thus withdrawn with a hydrogen containing gas, and returning catalyst thus separated and stripped directly to the lower portion of said elongated reaction zone at a substantially higher temperature than the gaseous mixture comprising hydrogen and carbon monoxide introduced into said reaction zone and at a temperature of at least 560° F.

9. The process of claim 8 in which the linear velocity of the gases passing upwardly through said elongated reaction zone is between about 8 and about 40 feet per second and the concentration of catalyst in the gases in said reaction zone is between about 3 and about 12 pounds per cubic foot of gas.

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