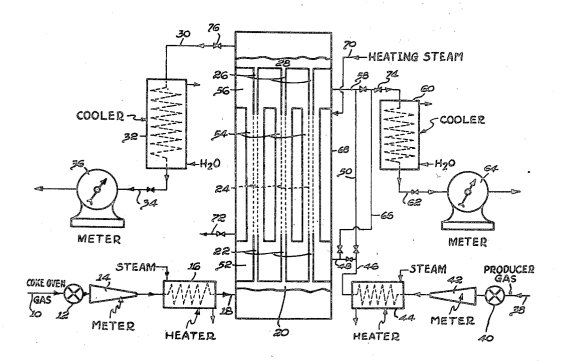
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METHOD OF CHANGING THE COMPOSITION OF A COMBUSTIBLE GAS BY DIFFUSION

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9 Claims. (Cl. 48—197)

This invention relates to gas diffusion for rearranging the composition of fuel gases. More particularly the invention relates to the revision of the composition of coke oven gas by diffusion when using producer gas as a sweep gas and 5 thereby revising both the coke oven gas and the producer gas.

The carbonization of coal in byproduct coke ovens produces a comparatively uniform quality of gas which, when the aromatic hydrocarbons 10 are removed, has a heating value of 530 to 580 B. t. u. and a comparatively low specific gravity. This gas is being widely used in cities for heating purposes. The construction of a coke oven battery requires that all of the ovens be operated 15 simultaneously so that the total or average output of gas is quite uniform and cannot be conveniently and economically varied to follow the daily or seasonal demand cycle.

it is customary to supplement the coke oven battery with water gas generators and then to mix the water gas with the coke oven gas to meet the demand. Some mixes of coke oven gas and water gas are not satisfactory because the B. t. u. 25 values may vary considerably and the gravity of the gas will also vary quite widely. The gas burning appliances do not operate satisfactorily with these variable types of gas so that it is imity of gas for city distribution. It has been found that when the specific gravity of the domestic gas is quite uniform, then the appliances will properly burn such gas.

The most desirable method of changing the 35 composition of coke oven gas is to vary the hydrogen content. When hydrogen is removed from the gas, the specific gravity and the B. t. u. value are increased. Therefore coke oven gas may be specific gravity more advantageously by diffusion to rearrange the constituents particularly the hydrogen content rather than attempting to obtain the desired gas by blending two or more different types of gases.

The primary object of the present invention is to provide a method of changing the composition of coke oven gas by selective separation of the hydrogen therefrom.

Another object of the invention is to provide 50 a method of separating hydrogen from coke oven gas by diffusion. In the diffusion method of hydrogen separation from coke oven gas, the hydrogen passes through a porous boundary as the gas flows in a continuous stream across one face 55 gas is at its maximum. Therefore, with coun-

of the boundary. The diffusion is substantially aided if another gas commonly termed "a sweep gas" is moved across the face of the boundary opposite the face across which the coke oven gas is flowing. The sweep gas is used to control the pressure distribution of the two gases on opposite sides of the boundary. The sweep gas has a flow through the boundary into the feed gas and also acts to sweep from the face of the boundary the hydrogen and other gases that have penetrated through the boundary from the feed gas side by diffusion.

The "pressure distribution" of the feed and sweep gases on opposite sides of a boundary involves several factors:

First, the size of the pores which determine the porosity of the boundary or diaphragm. For the revising of fuel gas, the boundary should have a pore size which is many times larger than the To take care of this variation in gas demand, 20 mean free path of the gas molecules. This mean free path of the molecules is the average distance which a molecule travels before colliding with another or an adjacent molecule. Depending upon the length and the desired resistance to flow of the gas through the pore, the width or diameter of the pore may be from 1 to 10,000 times the mean free path of the molecule.

Second, the pressure of the gas streams at opposite sides of the boundary or the average presportant to deliver a comparatively uniform qual- 30 sure differential across the pores of the boundary, or across the boundary.

Third, the area of the boundary, which takes into consideration the pressure differentials at the inlets or outlets of the boundary, or the average pressure differential over the entire area of the boundary. When the feed and sweep gas streams flow concurrently or in the same direction across the opposite faces of the boundary, the pressure differential at the entrance end is revised to obtain the desired heating value and 40 dependent on the pressure differential of the two inlet gas streams, and there will be a gradual decrease or increase of the pressure differential as the streams advance across the boundary area, depending upon whether the pressure drop due to flow resistance in the sweep stream is more or less than the pressure drop in the feed stream. If the feed and sweep gas streams flow in countercurrent direction along the faces of the boundary, then the pressure differential across the boundary at the entrance end of the sweep gas is at its maximum because the resistance to flow and the diffusion of the feed gas through the boundary has altered and lowered the pressure of the feed gas while the pressure of the sweep

tercurrent flow the diffusion rate will be the highest at the entrance end of the sweep gas, and lowest at the entrance end of the feed gas.

Fourth, the hydrodynamic flow of the sweep gas through the pores of the boundary. This 5 hydrodynamic flow is the free flow of sweep gas through the boundary pores not due to diffusion but due to the pressure differential across the boundary.

I have found that a low B. t. u. producer or 10 blast furnace gas which has a radically different composition than the composition of coke oven gas is an excellent sweep gas for facilitating the separation or redistribution of hydrogen in coke oven gas. Producer and blast furnace gases 15 have comparatively high specific gravities so that there is relatively small amount of diffusion of these gases through the boundary into the coke oven gas. On the other hand, the low specific gravity hydrogen readily passes through 20 the boundary and is swept away from the boundary by the high specific gravity gas. If hydrogen is passed through the boundary from coke oven gas into producer or blast furnace gas, the addition of hydrogen to these gases is very advantageous in that it lowers the specific gravity of the gases and substantially increases the B. t. u. heat values.

Accordingly, a further object of the invention is to utilize producer or blast furnace gas as a 30 sweep gas in the diffusion separation of hydrogen from coke oven gas.

In accordance with the relative volumes of sweep gas and feed gas being revised, it is desirable to maintain a fixed differential pressure 35 on opposite sides of the boundary, this differential pressure across the boundary being high if the boundary is only slightly porous and being comparatively low if the boundary is quite The temperature of the gases flowing across the boundary should be maintained above the dew points of any constituents in the gases and the velocity of the sweep gas should be such that it will sweep away from the face of the boundary the constituents diffusing through the boundary.

The pressure distribution of the sweep and feed gases with reference to the porous boundary of a diffusion apparatus depends upon the specific gravity of the gases, the pressure differential across the boundary, the velocity of movement of the gases across the face of the boundary and the area of the boundary. It is possible by the control of these various features to obtain a selective separation of hydrogen from coke

Typical compositions of coke oven, producer and blast furnace gases are shown in the following table.

TABLE I

Property	Producer	Coke Oven	Blast Fur-
	Gas	Gas	nace Gas
Specific Gravity B. t. u./Cu. Ft	0. 9	0. 4	0. 9
	130	586	85
Analysis: Carbon dioxide Illuminants	3. 5	1.8	11.5
Oxygen	0. 5	0. 2	27, 5
Carbon monoxide	30. 0	6. 3	
Hydrogen Nitrogen Methane	10.0 54.5 1.5	53. 0 3. 4 31. 6	1. 0 60. 0

My tests have shown that at the same time

gas may also be revised so that a large number of different types of gas may be produced. The revised producer or blast furnace gas which is enriched with the hydrogen and hydrocarbons of the coke oven gas is a good heating gas for coke ovens. Further, the revision of the sweep gas may be controlled to distribute the hydrogen and carbon monoxide in the ratio of molecular volumes of 1:1 or 2:1 to provide an excellent synthesis gas for the hydrogenation of carbon monoxide in the Fischer-Tropsch reaction.

A still further object of the invention is to provide a method of changing the composition of coke oven gas with a producer sweep gas to simultaneously upgrade the producer gas into a predetermined type of gas.

With these and other objects in view, the invention consists in the method of revising the composition of gas by diffusion as hereinafter described and particularly defined in the claims.

The various features of the invention are illustrated in the accompanying drawing which is a diagrammatic flow sheet of an apparatus in which the preferred method of revising coke oven gas by diffusion may be carried out.

Coke oven and producer gases are generally maintained at a comparatively low pressure (6 to 12 inches of water pressure). Accordingly the diffusion process for revising coke oven gas is carried out at substantially atmospheric pressure, that is, these gases will have a sufficient pressure to control their flow through the diffusion apparatus.

Referring to the drawing, the diffusion revision of coke oven gas may be carried out as follows:

Coke oven gas is introduced through a line 10 and flow regulator 12 into a meter 14 to establish a definite flow rate. The gas passes through a heater 16 which is preferably heated by steam to a temperature above the condensation temperature or dew point of any of the constituents in the gas at the boundary. From the heater 16 the gas passes through a line 18 into a chamber 20 45 of a diffusion apparatus. This gas is then distributed throughout the area of the chamber 20 and passes through tubes 22 which connect with porous diffusion boundaries 24 that form a continuation of the tubes 22. The streams of coke 50 oven gas pass through the diffusion boundary tubes, then through tubes 26 into a chamber 28 and thence flow through a line 30 into a cooler The cooler 32 is preferably cooled by water or other cooling medium in order to standardize the temperature of the gas so that it may flow through a line 34 through a meter 36 to measure its volume.

The producer sweep gas is introduced into the apparatus through a line 38 and flow regulator 40 to establish a definite predetermined flow rate of producer gas. The producer gas then passes through a meter 42 into a heater 44 where it is heated by steam to raise it to substantially the same temperature as the coke oven gas as maintained by the heater 16. This preheated gas then passes through a line 46 to a line 48 or a line 50 in accordance with the manner in which the sweep gas is circulated through the diffusion apparatus. If the sweep gas is to pass through 70 the diffusion apparatus in a parallel concurrent stream with the feed gas then the sweep gas will enter a chamber 52 of the diffusion apparatus through the line 48. The sweep gas will then be distributed throughout the chamber 52 and pass that the coke oven gas is being revised, the sweep 75 upwardly through the tubes 54 around the porous

boundary tubes 24. After passing across the porous boundaries, the sweep gas enters a chamber 56 and then flows out through a line 58 through a cooler 60 to be conditioned for passing through a line 62 into a meter 64. If the sweep gas is to pass across the porous boundaries in a stream parallel to the coke oven gas stream but countercurrent to the flow of the coke oven gas, then the sweep gas will pass through the line 50 to the line 58, thence into the chamber 56, then 10 down across the porous boundary tubes 24 into the chamber 52, and out through the line 48. From the line 48 the gas will pass through the line 66 to the line 58 and thence through the cooler 56 to the line 35 and thence through the 56.

50 and meter 64. To assist in obtaining a good temperature condition for diffusion, the tubes 54.

Sweep (producer gas) in= 643 C. F. H. (coke oven gas) out= 708 C. F. H. (coke oven gas) out= 708 C. F. H. (coke oven gas) out= 455 C. F. H. steam is introduced through an inlet 70 and water of condensation taken out through a line 12.

The pressure distribution across the porous boundary tubes 24 is maintained by means of a regulating valve 74 in the sweep gas line 58 and a regulating valve 76 in the feed gas line 30. The flow regulators 12 and 40 in the coke oven and producer gas lines respectively control the flow rate of coke oven gas and producer gas. The pressure of the gas flowing across the opposite sides of the porous barrier of the diffusion tubes is controlled by the valves 74 and 76. The valve 74 is controlled to maintain a pressure of the sweep gas on the outside of the tubes 24 slightly higher than the pressure of the feed gas within the tubes 24. This pressure differential then controls the hydrodynamic flow of sweep gas through the porous tubes and the diffusion of 35 constituents from one gas stream into the other gas stream. The pressure differential is small but is sufficient to maintain a hydrodynamic flow and definite diffusion through the diaphragm.

From Table I it will be seen that the hydrogen 4 content of coke oven gas is much higher than the hydrogen content of producer gas or blast furnace Hydrogen has a very much lower specific gravity than carbon dioxide, carbon monoxide, nitrogen or the hydrocarbons in coke oven gas. 4 Since the rate of diffusion through a porous diaphragm is roughly inversely proportional to the square root of the molecular weights of the gases. the hydrogen will have the highest velocity of any of the constituents in the feed and sweep gases. 5 Therefore, the revision of coke oven gas when using producer gas as a sweep gas will consist principally in the redistribution of hydrogen by diffusion through the porous boundary. By maintaining a higher pressure on the sweep gas side 55 of the boundary than on the feed gas side of the boundary, the free flow will be from the sweep gas into the feed gas. On the other hand, the coke oven gas constituents tend to set up a counterdiffusion through the boundary which is 60 opposed by the free flow and diffusion of the constituents of the producer gas through the porous boundary. The net result is an exchange of constituents passing through the boundary from each stream into the other. However, the higher 6 diffusion velocities of the lighter constituents in the coke oven gas, particularly hydrogen, will result in a net addition of hydrogen to the producer gas from the coke oven gas and a small redistribution of other constituents between the two gas 7 streams.

In Tables II to VII inclusive are shown data of the revision of a coke oven gas, of substantially the composition cited above, through a porous boundary in which tubes are used which have a 78

porosity of 6400 openings per square inch, the openings being substantially square with a dimension of 0.0035 inch on each side. These boundary tubes are made of a copper screen coated with corrosion resistant nickel. The data of Tables II to VII was obtained by using seven porous boundary tubes 11/2 inches inside diameter with an effective length of 24 inches. In all cases the gas was corrected to 30 inches of mercury pressure and 60° F. for the meter read-

TABLE II

۵۸		Sw	еер	Feed	
20		In	Out	In	Out
	H ₂ by Conductivity Cell. Orsat:	Per cent 10.5	Per cent 29.5	Per cent 51.7	Per cent 39.0
25	CO ₂ Illuminants	4.9 0.9	3. 2 0. 6	1.6 3.0	2.3
	O2	0.5	0.2	0.6	0.5
	H ₂	10.8	29.5	55, 6	40.1
	CH4C2H4	28. 5 1. 2	21. 9 5. 7	5. 6 29. 0	12. 5 21. 8
30	N ₂ (By difference)	53, 2	38.9	4.6	20.6
	Specific Gravity	0.870	0.697	0.370	0. 538

TABLE III

Feed (coke oven gas) in = 643 C. F. H. (coke oven gas) out = 816 C. F. H. Sweep (Producer Gas) in = 530 C. F. H. out = 357 C. F. H.

	Sweep		Feed	
	In	Out	In	Out
H ₂ by Conductivity Cell_ Orsat:	Per cent 10.5	Per cent 28.3	Per cent 51.7	Per cent 39. 2
CO ₂ Illuminants	4.9 0.9 0.5	3.7 0.7	1.6 3.0	2.6 2.2
H ₂ CO	10.8 28.5	1.0 30.1 21.1	0.6 55.6 5.6	1.7 39.8 12.8
CH ₄ N ₂ (By difference)	1. 2 53. 2	4. 5 38. 9	29.0 4.6	13. 4 27. 5
Specific Gravity	0.870	0.699	0.370	0. 570

TABLE IV

Feed (coke oven gas) in=187 C. F. H. (coke oven gas) out=307 C. F. H. Sweep (Producer Gas) in=476 C. F. H. out=356 C. F. H.

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		Sw	eep	Feed	
5		In	Out	In	Out
0	H ₂ by Conductivity Cell. Orsat: CO ₂	Per cent 10.5 4.0 0.0 0.8 11.8 28.7	Per cent 23. 1 3. 8 0. 0 1. 0 22. 1 24. 6	Per cent 54.3 1.7 2.8 1.5 53.9 6.7	Per cent 31.8 2.8 2.2 1.3 31.6 15.8
	CH4 C2H6 N2 (By difference)	0.0 1.5 53.2	2.0 1.8 44.7	17. 2 2. 9 13. 3	16. 1 0. 0 20. 2
	Specific Gravity	0.874	0.783	0.431	0.637

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TABLE V

Feed (coke oven gas) in=390 C. F. H. (coke oven gas) out=516 C. F. H. Sweep (Producer Gas) in=482 C. F. H. out=356 C. F. H.

					U
	Sweep		Fe	ed	
·	In	Out	In	Out	
H ₂ by Conductivity Cell.	Per cent 11.6	Per cent 23.8	Per cent 54.3	Per cent 39. 2	10
Orsat: CO ₂	4.9 0.0 0.2 12.4	4.0 0.0 0.1 24.0	1.7 2.6 0.7 54.9	2.7 1.9 0.7 39.7	
CO CH4 C ₂ H ₆ N ₂ (By difference)	28. 5 1. 4 1. 2 51. 4	23. 7 0. 1 1. 9 46. 2	6.3 24.6 0.6 8.6	13. 1 17. 4 1. 2 23. 3	15
Specific Gravity	0.889	0. 782	0.394	0. 560	

TABLE VI

	Sweep		Feed		
	In	Out	In	Out	
H ₂ by Conductivity Cell. Orsat: CO2	Per cent 11. 9 4. 9 0. 0 0. 2 12. 4 28. 5 1. 4 1. 2 51. 4	Per cent 34. 7 3. 8 0. 8 0. 5 34. 9 18. 4 8. 3 0. 4 32. 9	Per cent 55.7 1.7 2.6 0.7 54.9 6.3 24.6 0.6 8.6	Per cent 48. 1 2. 3 2. 5 0. 8 48. 9 8. 8 23. 0 0. 6 13. 1	
Specific Gravity	0.889	0.642	0.394	0.453	

TABLE VII

 $\begin{array}{lll} \mbox{Feed (coke oven gas)} & \mbox{in} = 584 \ \mbox{C. F. H.} \\ \mbox{(coke oven gas)} & \mbox{out} = 484 \ \mbox{C. F. H.} \\ \mbox{Sweep (Producer Gas)} & \mbox{in} = 400 \ \mbox{C. F. H.} \\ \mbox{out} = 500 \ \mbox{C. F. H.} \end{array}$

	Sweep		Feed	
	In	Out	In	Out
H ₂ by Conductivity Cell. Orsat: CO2	Per cent 11.0 4.9 0.0 0.2 12.4 28.5 1.4 1.2 51.4	Per cent 32. 2 4. 2 0. 8 0. 2 32. 3 18. 6 6. 7 0. 8 36. 4	Per cent 55. 7 1. 7 2. 6 0. 7 54. 9 6. 3 24. 6 0. 6 8. 6	Per cent 45.8 2.6 2.4 0.8 47.7 8.7 22.8 1.1
N ₂ (By difference) Specific Gravity	0.889	0.769	0.394	0.466

The data of Tables II, III, IV and V were taken at a time when the sweep gas was passed through the diffusion apparatus as illustrated in the drawings in a path concurrent with the flow of feed gas.

The data of Tables VI and VII were taken when the sweep gas flowed in a countercurrent path to the feed gas.

In Table VIII the data set forth in Tables II to VII inclusive is summarized to show the percentage gravity increase and the percentage B. t. u. decrease in the coke oven gas due to a single passage of the gas across the boundary. The percentage B. t. u. decrease is obtained by securing the B. t. u. value of the gas as based on the gas analysis wherein the B. t. u. value of

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illuminants and ethane is taken as 1600 B. t. u.'s per cubic foot, the carbon monoxide and hydrogen as 320 B. t. u.'s per cubic foot and methane as 1000 B. t. u.'s per cubic foot.

TABLE VIII

Coke oven gas

•	Percentage Gravity Increase	Percentage B. t. u. Decrease
Table II Table III Table IV Table IV Table VI Table VI Table VII	45. 4 55. 7 47. 8 42. 2 15. 0 18. 3	24. 5 36. 5 18. 9 20. 0 3. 67 2. 3

In Table IX is shown the percentage gravity decrease and the percentage B. t. u. increase for the producer gases revised by diffusion as shown in Tables II to VII inclusive. The B. t. u. value of the various gases are calculated on the basis described above with reference to Table VIII.

TABLE IX

Producer gas

	Percentage of Specific Gravity Decrease	Percentage of B. t. u. Increase
Table II	12, 2	50. 5 43. 5 30. 0 12. 5 60. 5 55. 5

When coke oven gas is used for domestic heating and cooking purposes, it is generally blended with coal gas or carbureted water gas. The spe-40 cific gravity of coke oven gas as it is produced from the ovens may vary from 0.38 to 0.45. It is very desirable to increase the gravity of coke oven gas in order to allow it to be blended or mixed with coal gas or carbureted water gas and burned 45 in the appliances usually used for carbureted water gas or coal gas. From Tables II to VII inclusive it will be seen that the revision of the coke oven gas by diffusion has raised the gravity from 0.4 to a specific gravity in the range of 0.453 50 to 0.637. The conditions set forth in Tables II and III are best adapted for conditioning coke oven gas to be blended with carbureted water gas for domestic purposes. The conditions set forth in Tables VI and VII are best adapted for 55 the revising of producer gas to make Fischer-Tropsch synthesis gas. When making coke oven gas to be blended with carbureted water gas the percentage gravity decrease of the producer gas varies from about 12 to 20% and the coke oven gravity is increased at the same time in the range of 42 to 50%. When making Fischer-Tropsch synthesis gas from the producer gas the gravity of the producer gas is reduced in the range of 25 to 28% and at the same time the specific gravity 65 of the coke oven gas is increased in the range of 15 to 19%.

From this data it will be seen that the composition of the coke oven gas has been quite widely changed, particularly with reference to 10 its hydrogen, and carbon monoxide contents. The specific gravity of the revised gases shown in the tables is very acceptable for domestic purposes. Further the B. t. u. value of the producer gas has been increased approximately 10%. It 76 will be seen also that the ratio of carbon monox-

ide to hydrogen in the reformed sweep gas may O be quite widely varied. In accordance with the data of Tables IV and V the molecular ratios of hydrogen to carbon monoxide in the revised producer gas are approximately 1:1, while in ac- Ω cordance with the data of Tables VI and VII the O molecular ratios of hydrogen to carbon monoxide in the revised producer gas are substantially 2:1. Such gases are well suited as synthesis gases for the Fischer-Tropsch process. Although nitrogen is present in these gases, it has been found that a high percentage of nitrogen is not detrimental to the catalytic hydrogenation of carbon monoxide with a catalyst in the Fischer-Tropsch reaction.

In the apparatus illustrated in the drawing, seven porous boundary tubes have been used. It is apparent, however, that the apparatus may be Q designed to use any desired number of tubes and the porous area may be varied in accordance with 20 the gases being treated in order to get the desired diffusion separation of products. In the diffusion operation there is always an exchange of gas from one side of the boundary into the gas at the fusion of the gaseous constituents may be controlled and modified in order to get a comparatively selective separation of the desired components.

The preferred form of the invention having 30 been thus described, what is claimed as new is:

1. A method of simultaneously changing the composition and specific gravity of coke oven gas and producer gas comprising: feeding coke oven gas as a stream along one side of a porous bound- 35 ary, passing producer gas in a stream to sweep across the opposite side of the boundary and controlling the pressure differential and the rate of flow of the gases on opposite sides of the boundary of a predetermined area to cause a 40 movement of hydrogen by diffusion through the boundary to the producer gas and nitrogen and carbon monoxide to the coke oven gas so that the specific gravity of the coke oven gas will be increased in the range of 42 to 50%.

2. The method defined in claim 1 in which the gases on opposite sides of the boundary are con-

ducted in concurrent paths.

3. The method defined in claim 1 in which a slightly higher pressure is maintained on the 50 rent paths. sweep gas side of the boundary than on the coke oven gas side of the boundary to cause a hydrodynamic flow of producer gas through the pores of the boundary.

4. The method defined in claim 1 in which the 55 file of this patent: composition of the coke oven and producer gases are brought to a finished composition in a single

pass across the boundary

5. The method defined in claim 1 in which the pore size of the boundary is in the range of one 60 to one hundred times larger than the mean free path of the coke oven gas molecules.

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6. A method of simultaneously changing the composition and specific gravity of coke oven gas and producer gas comprising: feeding coke oven gas as a stream along one side of a porous boundary, passing producer gas as a stream to sweep across the opposite side of the boundary, controlling the pressure differential and the rate of flow of the gases on the opposite sides of the boundary of a predetermined area to increase by 10 diffusion the hydrogen content of the producer gas and the carbon monoxide and nitrogen content of the coke oven gas while increasing the specific gravity of the coke oven gas in the range of 42 to 50%.

7. A method of simultaneously changing the composition and specific gravity of coke oven gas and producer gas comprising: feeding coke oven gas as a stream along one side of a porous boundary, passing producer gas as a stream to sweep across the opposite side of the boundary, controlling the pressure differential and the rate of flow of the gases on the opposite sides of the boundary of a predetermined area to increase by diffusion the hydrogen content of the producer gas and the other side of the boundary but the rates of dif- 25 carbon monoxide and nitrogen content of the coke oven gas while decreasing the specific gravity of the producer gas in the range of 12

to 20%.

8. A method of increasing the specific gravity of coke oven gas while converting a producer gas into a Fischer-Tropsch synthesis gas comprising: feeding a coke oven gas having a gravity of approximately 0.4 as a stream along one side of a porous boundary, passing a producer gas having a specific gravity of approximately 0.9 along the opposite side of the boundary as a sweep gas, controlling the rate of flow and pressure differential of coke oven gas to producer gas on opposite sides of a boundary of a predetermined area to increase the hydrogen content of the producer gas by diffusion in the range of two volumes of hydrogen to one volume of carbon monoxide while simultaneously increasing the specific gravity of the coke oven gas and decreasing the specific gravity of the producer gas in the range of 25 to 28%.

9. The method defined in claim 8 in which the coke oven and producer gases are caused to flow on opposite sides of the boundary in countercur-

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