UNITED STATES PATENT OFFICE.

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GILSONITIC PRODUCTS.

Original application aled August 28, 1919, Serial No. 320,514. Divided and this application aled April 3, 1922. Serial No. 549,186.

10 new and useful Improvements in Gilsonitic tures, unless otherwise specially noted.) cation.

15 cially to products and materials obtainable from the still begins at a temperature of ment. We have found that there can be beyond coking temperatures. obtained from gilsonite and gilsonitic sub- Up to the point where the vapor tempera- 70 stances semi-drying oils characterized by ture approaches some 550° F., the distilla-20 various useful properties.

plication, Serial No. 320,514, filed August heat. At this point, however, some exother-

their manufacture".

The various phases of our invention will become excessively rapid. Unless, therefore, be specifically described hereinafter in ac- the previous heating has been especially cordance with the best mode of applying slow, it is necessary to reduce the application them in a practical way at present known of heating very greatly as this critical point 80 to us.

As a gilsonitic material to be treated, we liquid may not be carried over into the conordinarily prefer gilsonite distillate to na- denser, and clog it up so as to render it untive gilsonite itself. We have obtained a usable. In practice, it will usually be found much greater yield of sulphonic products convenient to cut down the fire some 100° F., 85 from the treatment by preparing the dis- in advance of this temperature. Once this 35 tillate sometime in advance and allowing it critical point is well passed, the fire may be to "age" for a considerable period before increased and the distillation pushed on as being treated. Such distillate may be pre- rapidly as desired to its conclusion. In

pared as follows:

charged into an iron or steel still, fired with measured on the solid coke product collectgas or oil (or otherwise suitably heated), ing in the bottom of the still. and connected to a suitable condenser,— While the vapor coming off from the still such as an iron pipe condenser water-cooled. may all be led into the iron pipe condenser, 95 Any convenient quantity of gilsonite may be as above suggested, and all of the strictly charged,—say 600 pounds to several tons. vaporous portion thereof there condensed The still being closed and heat applied and collected as a liquid, a slightly different gradually, the gilsonite will liquefy and col- procedure is preferable. A minor portion of lect in the bottom of the still, and vapor and the vapor is condensable at relatively high 100 gas evolving from it will fill the top of the temperatures; and at the temperature in the still and pass over into the condenser. As condenser, the corresponding condensate is the heating progresses, temperature readings of such thick, gummy consistency that it should be taken from time to time on the would tend to clog it up. It is desirable,

To all whom it may concern: body of vapor in the upper portion of the Be it known that we, Charles N. Forrest, still, as well as on the body of liquid beof Rahway, in the county of Union and State neath it. (Excepting as there may be spe- 55 of New Jersey, Harold P. Hayden, of cial occasion for distinguishing them, we 3 Raritan Township, in the county of Middle- here and hereinafter comprehend mingled sex and State of New Jersey, and Orin R. gases and vapors under the single term Douthett, of Beaver Falls, in the county of "vapor", for the sake of brevity. For con-Beaver and State of Pennsylvania, citizens venience, we refer to various stages of opera- 60 of the United States, have invented certain tions by the corresponding vapor tempera-Products, whereof the following is a specifi- The effect of the progressive heating of the gilsonite is to gradually break it up and de-Our invention relates to gilsonitic prod- compose it chemically,—whence, mainly, the 65 ucts and their manufacture, and more espe- evolution of vapor. The coming off of vapor from gilsonite through sulphonation treat- some 165° F., and continues up to or even

tion may be carried out as rapidly as the This application is a division of our ap- contents of the still can be made to absorb 28, 1919, entitled "Gilsonitic products and mic or other peculiar action occurs, so that 75 the evolution of vapor in the still tends to is approached, in order that undistilled practice, the heating need not be carried 90 Gilsonite as received from the mine is further than a temperature of some 850° F.,

normally liquid products be condensed and coverable by a similar operation in conneccollected separately, without entering the tion with conversion thereof into fuel gas condenser at all. This can advantageously 5 be taken care of by making the pipe leading gas-producer. The water eliminated from 70 from the still to the condenser of substantial the crude distillate also contains ammonia length and providing it with a trap. With this arrangement, the heavier portion of the normally liquid products will be condensed 10 by the cooling effect of the atmosphere upon the pipe (which thus acts as a sort of auxiliary condenser), and will collect in the trap, whence it can be drawn off from time to time and added to the liquid products drawn 15 from the condenser itself. The trap also affords a measure of protection against the effects of too rapid heating at the critical temperature above mentioned.

The crude distillate thus obtained contains 20 some 2 to 5% water, which may be eliminated by settling it out in a settling tank.

We prefer not to treat the water-freed primary, crude distillate thus produced directly, in gross, but to separate it into a 25 plurality of different portions by fractionation or reduction. The following examples (wherein the temperatures given are the vapor temperatures in the still, unless otherwise stated) will sufficiently illustrate the or more times. most convenient methods of redistillation:

(1) Condense separately the vapors coming from the still up to 475° F. and from jecting the purified gilsonitic oil to the ac-475° to 600° F.,—subsequently drawing off as residuum the oil unvaporized at the latter phonating reagents which we prefer to use 35 temperature, or allowing it to remain and mix with the next charge of crude oil.

(2) Condense separately the vapors up to 475° F.: from 475° F. to 650° F.; and from 650° F. until the temperature of the mate-40 rial in the bottom of the still is about 850° F. By this procedure, an amount of coke equal to about 2% of the crude oil charge will be produced.

In either of these cases, considerable de-. 45 structive action occurs in the production of the higher fractions,—especially the last.

(3) Three fractions nearly similar to those described under (2) may be obtained at temperatures some 100° F. lower than those mentioned by carrying out the redistillation as described in U.S. Patent 877,620, granted January 28, 1908, to Wells, blowing carbon dioxide or other inert permanent gas through the liquid in the still. In this case, under (1).

The gas from the primary distillation contains ammonia, which can be segregated as ammonium sulphate by a sulphuric scrub-65 bing operation. The coke from both distil-

therefore, that this heavier portion of the lations also has a nitrogeneous content reby incomplete combustion in a by-product in solution, which may, if desired be also converted into ammonium sulphate. For these operations for recovery of ammonia may be used the unconsumed portion of the 75 sulphuric reagent employed in the purification and sulphonation treatments hereinafter described.

Of the distillates thus produced, we generally prefer to treat the intermediate one. 80 Prior to sulphonation treatment, however, we prefer to subject the gilsonitic material to preliminary purifying treatment with a sulphuric reagent, thus removing therefrom organic bases, olefines, and other impurities 85 that might alter or impair the character and utility of the final products. This preliminary sulphuric treatment may be carried out by agitation of the gilsonitic material with about 5% of sulphuric acid of oil of 90 vitriol grade,—i. e., about 66° Bé.,—for about one half hour. It is desirable to repeat the operation (using fresh acid) one

After this preliminary step, "sulphona- 95 tion" treatment may be carried out by subtion of a sulphonating reagent. The sulare oil of vitriol, such as mentioned above, 100 and fuming sulphuric acid or "oleum." It is desirable to conduct this treatment in a cast iron vessel, using mechanical agitation rather than air agitation in order to avoid such contamination with atmospheric mois- 105 ture as might interfere with the desired re-

action. We have found that the quality of the final products is improved by carrying out this treatment progressively, with successive 110 portions of reagent, and segregating the reaction products, as formed, from the material being treated, so that each of the successively acting portions of reagent shall be perfectly fresh and untainted with such re- 115 action products. Further improvement can be obtained by starting with relatively weak reagent, and using more and more reactive or energetic reagent as the treatment pro-55 the destructive action is much less. It is gresses. This may be done in a succession 120 advantageous to pass the gas and vapor of separate operations upon a single lot of coming off through a filter of fuller's earth material with different portions of reagent, in the dome of the still, on its way to the rather than in one single operation,—say condenser. A residuum of heavy oil will four such treatments, each with an amount 60 preferably be left in the still, as described of reagent equal to 20% of the material 125 treated. Thus for the first treatment, the reagent may be oil of vitriol alone; for the second, a mixture of 15 parts oil of vitriol and 5 parts oleum; for the third, a mixture of oil of vitriol and oleum in equal parts; 130

material.

same purpose. This gilsonitic sulphonation absent). residual oil as thus purified is a thin, clear Coming, now, to the further disposition oil: substantially colorless, orderless, and or treatment of these separation products.— 35 tasteless; and suitable for about the same the dilute acid of the bottom layer may be 100 uses as highly refined petroleum. If de- used for the recovery of ammonia as above sired, the distillation above mentioned may described. The "top layer" (when present) be carried out fractionally, so as to yield may be neutralized with sodium hydrate particular purposes.

tained from different sulphonation treat- slow-drying or "semi-drying" character; ments be further treated or worked up in- and usually of dark, reddish-brown color dividually, since the various treatments yield and slight, inoffensive odor, somewhat sugfinal products having somewhat different gesting that of vaseline. It is soluble in 110 sulphonating reagent, water-soluble organic after sufficiently prolonged exposure to the the mother oil by chemical action incident as a semi-drying oil; but when mixed with 115 to the sulphonation treatment. It also con- boiled linseed oil, it so modifies the drying tains, as impurity, an inconsiderable amount properties of the latter that it dries homosequently segregated, be neutralized along surface film. It also seems to render the 55 with that obtained as above described), and fully dried linseed oil tougher and more 120 some small amount of other organic prod- elastic. ucts. The proportions of the sulphonic and We have hereinbefore spoken of sulphoother reaction products in different lots of nating reagents, sulphonation treatment, product, in particular, may be entirely ab- products, etc., using these expression in ref- 125 sent from the sludge of the last sulphonation erence to the chemical character and mode treatments. The condition of the materials of use of the reagent employed in the prin-

We have found that separation of the ing therefrom. We do not, however, mean 130

for the fourth, a mixture of 5 parts oil of sulphonic material mentioned above fron vitriol and 15 parts oleum. Each of these the rest of the sludge can be effected by adtreatments may last about two hours. dition thereto of a solvent of the sulphonat-If, on the other hand, it is desired to carry ing reagent: for this purpose, the sludge out sulphonation treatment all at once, in a may be diluted with an equal weight of 70 single operation, rather than progressively, water. Preferably the water is placed in it may be done by mechanical agitation for a lead-lined, water-jacketed vessel, and the some eight hours,—using, say, an amount of sludge gradually introduced at such a rate sulphonating reagent equal to 80% of the that the temperature shall not rise exces-10 material treated and consisting of 50 parts sively,—110° F., being a perfectly safe 75 oil of vitriol and 30 parts oleum. limit. The operation is decidedly ex-At the end of each treatment or operation, othermic, and sulphur dioxide is copiously the charge in the agitator is allowed to stand evolved. The dilute sludge is allowed to an hour or more, when it will be found to stand some eight to twelve hours,—more have separated into two distinct layers; a or less,—at a temperature of 150°F.. or 80 top layer of residual or mother oil, and a thereabout. During this period, separation bottom layer of acid sludge. This latter into three distinct layers takes place, and should be drawn off and thus segregated these may be successively drawn off and thus prior to the next treatment of the mother segregated. The bottom layer is a dilute aqueous solution of the excess of sulphonat- 85 The residual oil remaining after the last ingreagent, discolored by some slight amount treatment may be neutralized, as with of organic impurity; the middle layer comsodium hydrate solution (preferably of prises the water-soluble organic sulphonic specific gravity greater than 30° Bé., in material, contaminated with an inconsiderorder to avoid emulsification). After the able amount of the solution that forms the 90 excess of caustic has been drawn off, the re- bottom layer and with a small portion of the sultant alkaline oil should preferably not ingredients of the top layer; and the top be washed, on account of its emulsifying layer contains the reaction oil component tendency; but it may instead be distilled and the residual mother oil impurity, etc., with steam, in order to improve its color, mentioned above. (In some cases, as indiand filtered through fuller's earth for the cated above, the "top layer" may be entirely

different oils more especially suitable for and washed with water until free from alkali. The resulting gilsonitic sulpho- 105 It is preferable that the lots of sludge ob- nation sludge oil is a viscous liquid; of properties. This sulphonation sludge con- benzole or naphtha and insoluble in alcotains, as principal components, unconsumed hol, when freshly prepared; and vice-versa sulphonic products, and oil produced from air. Not only is it adapted for general use of the mother oil itself (which may, if sub-geneously, without formation of the usual

sludge vary considerably; the oil reaction and sulphonation reaction or sulphonic in the sludge is that of solution in the uncon-cipal treatment described above, or to sumed sulphonating reagent. the useful properties of products result-

sense of having sulphonic radicals,—since we are without clear evidence on this point.

Having thus described our invention, we

claim:

1. A semi-drying, viscous, gilsonitic sul-

phonation sludge oil.

2. A gilsonitic oil; viscous and slow-dryin ing; soluble in benzole and naphtha and insoluble in alcohol when fresh, and vice versa when dried.

. 3. A semi-drying, viscous gilsonitic oil;

by this language that our sulphonic prod- having the property, in admixture, of causucts are true sulphonic substances in the ing boiled linseed oil to dry homogeneously 1 instead of with a surface film.

In testimony whereof, we have hereunto signed our names at Maurer, N. J., this 29th

day of March 1922.

CHARLES N. FORREST. HAROLD P. HAYDEN. ORIN R. DOUTHETT.

Witnesses:

D. J. Desmond,