EFFECT OF FRACTAL DIMENSION OF ASPHALTEN AGGREGATES
ON MACRO PROPERTIES OF NATIVE OIL
Aleksandr A. Zlobin
Perm National Research Polytechnic University (29 Komsomolskiy av., Perm, 614990, Russian Federation)

The paper discusses the complex analysis of the energy and structural features of the structure of asphaltenes of nanoggregates that was experimentally conducted using the data of the nuclear magnetic resonance method. Fractal dimension of the core of complex structural units (CSU) for two alternative groups of native oil was calculated, which is equal to 2.040 and 1.556 units. It is established that crude oils with a low fractal dimension of nucleus have a 1.5-5.0 times higher dynamic viscosity. Theoretical substantiation of the physical phenomenon was carried out. It is established that oils of low fractal dimension due to their structural individuality of the structure of the paramagnetic framework have an increased natural inhibitory capacity of the asphaltenes of the nucleus in relation to growth and fallout of solid paraffins forming organic was deposition. According to temperature studies simulating cooling of the flow of oil during lifting in the tubing, it was established that the growth of the radius of the CSU nucleus during aggregation of asphaltenes is well described by the linear model. The aggregation intensity is 3.6 times higher in oil with more than low fractal dimension. In addition, it was revealed that the coefficient of temperature aggregation of asphaltenes depends on the radius of the initial material, and for the oil with a high fractal dimension it is described by a monotonous logarithmic model. There was an abnormal nonlinear aggregation established for low-dimensional oil at small radii of the initial material, which leads to a violation of the monotonic nature of the change in the coefficient of aggregation. According to results of experiments that simulate the deposition of paraffins on a metal surface, it has been established that the effectiveness of wax inhibitors in oil with a low fractal dimension is 5–49 % higher compared to a group of high-dimensional oil. The results obtained in the work can be used in practice when developing modern methods of managing the properties of petroleum dispersed systems in the development, production and processing of hydrocarbons.

Key words: dimensions of asphaltenic core, structure of complex structural units, fractal dimension of nucleus, temperature aggregation of asphaltenes, effectiveness of wax inhibitors.

ВЛИЯНИЕ ФРАКТАЛЬНОЙ РАЗМЕРНОСТИ АСФАЛЬТЕНОВЫХ АГРЕГАТОВ НА МАКРОСВОЙСТВА НАТИВНОЙ НЕФТИ
А.А. Злобин
Пермский национальный исследовательский политехнический университет
(614990, Россия, г. Пермь, Комсомольский пр., 29)

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Ключевые слова: размеры асфальтенового ядра, строение сложных структурных единиц, фрактальная размерность ядра, температурная агрегация асфальтенов, эффективность ингибиторов АСПО.

В представленной работе с использованием данных метода ядерно-магнитного резонанса экспериментально проведен комплексный анализ энергетических и структурных особенностей строения асфальтеновых наногрегатов и рассчитана фрактальная размерность ядра сложных структурных единиц (ССЕ) для двух альтернативных групп нативной нефти, которая составляет 2,040 и 1,556 ед. Установлено, что нефть с низкой фрактальной размерностью ядра обладает более высокой (в 1,5–5,0 раз) динамической вязкостью, и проведено теоретическое обоснование данного физического явления. Выявлено, что нефть с низкой фрактальной размерностью в силу своей структурной индивидуальности строения парамагнитного каркаса обладает повышенной естественной ингибиторной способностью асфальтенов ядра по отношению к росту и выпадению твердых парафинов, формирующих органические асфальтеносмолопарафиновые отложения (АСПО). По данным температурных исследований, моделирующих охлаждение потока нефти при подъеме в насосно-компрессорных трубах, установлено, что динамика роста радиуса ядра ССЕ при агрегации асфальтеновых частиц хорошо описывается линейной моделью, при этом интенсивность агрегации выше в среднем в 3,6 раз в нефти с более низкой фрактальной размерностью. Дополнительно выявлено, что коэффициент температурной агрегации асфальтенов зависит от радиуса начальной затравки и для нефти с высокой фрактальной размерностью описывается монотонной логарифмической моделью. Для нефти с низкой размерностью установлена аномальная ненормальная агрегация при малых радиусах начальной затравки, что приводит к нарушению монотонного характера изменения коэффициента агрегации. По результатам опытов, моделирующих осаждение парафинов на металлической поверхности, установлено, что эффективность действия ингибиторов АСПО в нефти с низкой фрактальной размерностью на 5–49 % выше по сравнению с группой высокоразмерной нефти. Полученные в работе результаты могут быть использованы на практике при создании современных методов управления свойствами нефтяных дисперсных систем при разработке, добыче и переработке углеводородов.

Alekseandr A. Zlobin (Author ID in Scopus: 36712914500) – PhD in Engineering, Associate Professor at the Department of Oil and Gas Technologies (tel.: +007 919 457 05 38, e-mail: ZlobinAA55@gmail.com).

Злобин Александр Аркадьевич – кандидат технических наук, доцент кафедры нефтегазовых технологий (тел.: +007 919 457 05 38, e-mail: ZlobinAA55@gmail.com).
Introduction

Development and implementation of modern nanotechnologies is one of the priority areas for increasing the efficiency of recovery technologies and processing of crude hydrocarbon.

Today, there is no uniform standard that could describe what the nanotechnologies and nano goods are. On the territory of the Russian Federation the meaning of nanotechnology is established in GOST R 55416-2013 [1] (identical to the international document ISO/TS 80004-1: 2010 [2]) “Nanotechnology. Part 1. Basic terms and definitions”, in particular: a set of technological methods used to study, design and manufacture materials, devices and systems, including control and management of the structure, chemical composition and interaction of constituent elements of the nanoscale on purpose. The practical aspect of nanotechnologies includes production of devices and their components necessary for creation, processing and manipulation of atoms, molecules and nanoparticles. In a broader sense, this term also covers methods of diagnostics, characterology, and studies of such nanoobjects.

One of the most important theoretical and practical questions facing nanotechnology today is how to get molecules to group themselves in a certain way, to organize themselves and eventually obtain materials and objects with new physical and chemical properties [3]. That opens up opportunities for more efficient management of dispersed systems (oil, emulsions, drilling muds, surfactants, lubricating oils, additives etc.) in various oil and gas production processes [4-6], which makes nanotechnologies to be relevant.

In the present paper that continues the cycle of works on nanoobects a complex analysis of two groups of native oil was carried out. The groups differ in the magnitude of the fractal dimension of asphaltene aggregates. The effect of fractal dimension on various technological macroproperties of produced hydrocarbons is studied as well.

Aggregation processes and fractal dimension of asphaltene complexes

For the moment, physical and chemical properties, phase states of asphaltene aggregates and their dependence on the critical parameters of the system are quite well known. Nevertheless, it is emphasized that “current knowledge is sufficient for forecasting, but still insufficient to control the phase behavior of asphaltenes and oil systems as a whole” [13, 14].

Processes of self-organization of disperse systems in the general case are rather complicated and not completely understood. Thus, in a theory of water disperse systems the main role in aggregation is played by the charge Coulomb interactions of particles and potentials of charged interphase surfaces, which are well described by the Deryagin-Landau-Fervey-Overbeck theory [15-17].

Everything is much more complicated in oleo-dispersed systems. Firstly, there are no Coulomb forces there. Secondly, the basic van der Waals interactions (orientational, induction and dispersion) have specific quantum properties and depend on the ratio of polar and nonpolar molecules. Thirdly, particles of a dispersed phase, that are also called complex structural units, surrounded by molecules of the dispersion medium, have a complex composition of structural elements such as a core, outer and inner solvate shells (Fig. 1), which can be transformed and cardinaly rearranged under external impact of physical and chemical fields. For example, under the influence of temperature asphaltene core and solvate shell continuously change dimensions; in each oil it happens according to its own law [11]. An increase in radius of a CSU complex characterizes the aggregation process; decrease characterizes the peptization of particles.

Processes of aggregation of supramolecular asphaltene structures consist in convergence due to Brownian motion and collision of individual active...
complexes, which leads to the formation of a larger asphaltene core and CSU as a whole. In turn, peptization processes, inverse aggregations, consist in the spontaneous thermodynamically favorable dispersal (fragmentation) of a nucleus and appearance of a large number of smaller complexes of CSU. Any disperse system, which includes oil, tends to get a dynamic equilibrium between aggregation and peptization processes, which causes the kinetic stability of the nanostructure of aggregates under the influence of the temperature factor $KT$ ($T$ – temperature, $K$ – constant L.E. Boltzmann).

The state, structure of CSU and self-assembly processes of the asphaltene core are determined theoretically by energy balance of intermolecular interactions between the dispersion medium and dispersed phase [18, 19], taking into account the permeability of the solvate shell made of resins. It was established in [8] that in case of oleo-dispersed systems the relative activation energy can be used as a parameter for aggregation of asphaltenes in oil $\frac{X(E_c)}{E_c}$, that characterizes the resulting driving force in the self-assembly of CSU in oil. Using this approach, a graphical function of the aggregation of the CSU nucleus in native oil was constructed (Fig. 2), taking into account the real content of hydrocarbons and asphaltene and resin substances.

Data analysis shows that with a significant difference in energy of the dispersion medium and dispersed phase, size of nucleus on the left and right tends to an equilibrium state with a minimum radius of 10 nm (Fig. 2), which confirms the theory of extreme states of CSU [18]. Energy of the medium (solvent) and dispersed phase is the limiting condition for extreme metastable state of the nucleus, to which both branches of the graph tend but do not really reach it. The obtained “resonant” type of the graph in Fig. 2 is universal for both oleophobic and oleophilic systems, which is confirmed by the data of [20], where the mechanism of growth of particles of water metal sol. But pH value of the medium is a driving force of aggregation in [20], which regulates the potential of the electrostatic interaction of initial particles.

The function of change of CSU nucleus size from the aggregation parameter for native oil in Fig. 2 consists of two branches such as left and right which characterize the different energy potential of a nucleus. The left branch describes the oil in which $X(E_c) < 1$, i.e. the energy of a dispersion medium is less than energy at the interphase boundary with the nucleus. The right branch in Fig. 1 describes an oil group of condition $X(E_c) > 1$ and, respectively, the energy of a dispersion medium is greater than the energy of a dispersed phase. In general, the kernel aggregation function in Fig. 2 is asymmetric, which indicates a significant difference between these two groups of oil. Using the analogy with aggregation in water colloids, we can presumably speak about existance in the oil of the fast mechanism [21] (left branch) and slow (right branch) aggregation of particles.
It was established that aggregation of asphaltenes in native oil occurs according to two different scenarios that are caused by individual structural and energy properties of the CSU.

For the analysis oil with left-hand aggregation is referred to the first experimental group with a “strong core” and “weak core” for oil with right-hand aggregation.

Strong nucleus is characterised by a step-by-step increase in the radius of the aggregate (see Fig. 2). Weak nucleus is characterised by smoothly and monotonously rearrangement over a wider range of aggregation parameter change $X(E_c)$ (0.45 vs. 0.25 fraction units).

The complex analysis carried out comparing the properties of two groups of oil that are not equivalent in terms of aggregation showed that for identical core radii, the following differences are observed. Average oil density in the first group is somewhat larger than in the second group. Average thickness of the solvate shell of resins in the first group is greater than in the second one by 32-35 %. The radius of the CSU complex in the first group is greater than in the second one by 18-20 %. The sulfur content in the first group is slightly higher than of the second group.

Regardless all features mentioned, different energy activity of the CSU nucleus is the main difference between the groups. That is clearly seen from the data of the experiment in Fig. 3, where, firstly, the average activation energy [22] at the boundary with the dispersed phase in the first group is higher than in the second one by 47.8 % rel. (18.96 against 12.83 kJ/mol). Secondly, the change in the activation energy differs significantly in the groups. Thus, for the second group, increase in the nuclear radius is accompanied by an almost linear increase, and then a decrease in the activation energy after reaching an extreme size of 45 nm.

There is a nonlinear (oscillation) character of the activation parameter observed caused by at least three extremal states separated by equidistant intervals of 10-12 nm.

Thus, the first group has critical sizes of nucleus, under which self-consistent amplification or attenuation of the activation potential occurs. Such features do not appear in the second group or they are rather weak in magnitude.

Extremes of native oil observed in the graphs of Fig. 3 correspond to metastable states of conversion of the CSU structure with change in the content of the dispersed phase, which is confirmed by the data in Fig. 4 that shows the experimental graphs of the relationship between the thickness of the solvation shell and CSU radius. It is found that the thickness of the adsorption shell for the second group (chart 2) remains the same at 24 ± 2 nm up to the boundary radius and then increases slightly, which characterizes a weak dependence of solvate shell thickness on nuclear radius. In other words, a core with low activity is surrounded by a relatively homogeneous thin solvate shell. Comparing the graphs in Fig. 3 and 4 it is clear that activation energy at the boundary with nucleus is negatively related to the ratio of solvate shell thickness to the radius of the nucleus $h_{ads}/R$, i.e. core shielding factor.
At the same time, there is a nonmonotonic process of extreme changes in thickness of a solvate shell observed for the first group with a strong nucleus (see Fig. 4, graph 1) as a function of geometric dimensions of the nucleus with a tendency to increase as the size of the CSU increases. In this case activation energy does not correlate with the screening factor like in the second group.

The structural differences of the CSU complexes in the groups established above suggest that the main reason is the core itself as the central potential force. That force affects all peripheral objects and exchange processes between the nucleus and dispersion medium.

It is assumed that all the CSU nuclei are spherical and have a uniform porosity, since they consist of individual small enough (about 2-2.4 nm) compact molecules of asphaltenes [23], but this is not always so. Therefore, in order to study the internal structure of the nucleus itself, representations of the fractal geometry of nanoobjects were used [24, 25].

Today, fractal models of formation of asphaltene nanoaggregates are fruitfully used to describe the processes of self-organization of oil dispersed systems [26, 27]. According to those processes, the asphaltene aggregate is a paramagnetic frame of fractal geometry [28]. The main feature is that the radial density function \( \rho(R) \) of the fractal aggregate is variable and rapidly decreases in power-law dependence with increasing distance from the center of the aggregate \( R \):

\[
\rho(r) = \rho(0)(R/r)^{D-d},
\]

where \( \rho(0) \) – initial density in the center of the unit; \( r \) – radius of particles from which the fractal aggregate is assembled; \( D \) – fractal (fractional, noninteger) dimension; \( d \) – dimensions of space (for a sphere \( d = 3 \)).

Fractal dimension reflects a special property of the aggregate, in particular the maximum level how it could fill the space for a fixed mass. The smaller the fractal dimension of \( D \), the more friable (with a smaller density of dendritic branches) the volumetric nanostructure arises in space and the greater the porosity of the aggregate [9].

Real density of the asphaltene core of native oil sample was estimated using the CSU structural elements obtained from the experiment. If the density of oil is known \( \rho_o \), and partial densities of the dispersion medium \( \rho_{d.m.} \), and density of resins \( \rho_{res} \), then it is possible to estimate the average density of asphaltenes in oil in situ without the deposition of asphaltenes by hexane or n-heptane [10].

The Fig. 5a shows average density of the asphaltene core of CSU plotted as a function of the radius for two groups of native oil of different deposits. It is established that as the radius of asphaltene aggregates increases, the density decreases by more than a factor of 2 in terms of the power law, which confirms experimentally the fractal nature of the nucleus of native oil.

However, due to the influence of various statistical factors, the obtained data on the calculated additive three-parameter model in Fig. 5a do not allow to estimate the fractal dimension in each oil group. More accurate methods are used for that. For example, a mass equation is frequently used that relates the number of particles \( N \) in the aggregate and their mass from the radius of the aggregate [29, 30]:

\[
N(r) = k(R/r)^D.
\]

Using the technique, linear graphs of interconnection between the mass of asphaltene nanoaggregates and nuclear radius for two established oil groups are obtained in double logarithmic coordinates and given in Fig. 5b.

That is seen from the Fig. 5b that mass method gives a clear differentiation of the sample in two groups. That also allows to calculate the fractal dimension. The experimental dependences obtained from (2) are described by power equations

1\textsuperscript{st} group – \( Y = 1.4519 \cdot 10^{-3}X^{1.556}, \rho = 0.960 \);  
2\textsuperscript{nd} group – \( Y = 2.857 \cdot 10^{-3}X^{2.040}, \rho = 0.985 \).

It follows that the fractal dimension of the asphaltene core for the oil of the first group is \( D = 1.556 \), respectively for the second \(-2.040 \), and deviation \( \Delta D \) between them is 31.1 %. Consequently, the first oil group is characterized by a lower, and the second, respectively, high fractal dimension of the asphaltene core.

The results obtained confirmed that the “strong” core, in comparison with the “weak” one, differs by the more complex volume structure of CSU fractal framework. Comparing the
Fig. 5. Dependence of average density of the CSU nucleus (a) and interconnection of
the mass of asphaltene particles (b) with radius for native oil samples. Code – oil groups

Effect of fractal dimensions
on macro-properties of oil

Dynamic viscosity. Oil is a complex dispersed system. If the oil contains asphaltene and resin
matter then various features of viscous flow appear [32]. The established fractal features
of the structure of the CSU nucleus of two groups of oil have a direct effect on the
technological macroparameters of native oil, in particular, the dynamic viscosity. It is known
that oil viscosity changes if fractality is considered [33].

Analysis of dependence of oil dynamic viscosity on content of asphaltenes, given in Fig. 6,
shows that having the content of asphaltenes changed oil viscosity in groups obeys following
equations:

\[
1^{\text{st}} \text{ group: } \ln Y = 0.816X + 1.083, \rho = 0.950, \\
\sigma = 0.182; \tag{5}
\]

\[
2^{\text{nd}} \text{ group: } \ln Y = 0.516X + 0.895, \rho = 0.981, \\
\sigma = 0.108. \tag{6}
\]

It can be seen from Eqs. (5) and (6) that experimental data for two groups are well described by a linear model, but the
following differences appear. The statistical parameters given in (5) and (6) characterizing the
tightness of the bond (dispersion \(\sigma\) and correlation coefficient \(\rho\)) show that there is a
practically functional dependence of the viscosity on the content of asphaltenes for the second oil
group. In addition, if we do not include the explicitly fallen off point in Fig. 6, then \(\rho\) is
additionally reduced in 5 times and approaches zero, and the correlation coefficient \(\rho\) becomes
equal to one in this case.
At the same time, the first group is characterized by a purely statistical relationship, which is characterized by a higher (1.7 times) variance and low correlation coefficient, which indicates the effect of additional statistically significant parameters, in addition to the content of asphaltenes. For example, if the content of asphaltenes is normalized to the radius of the core, i.e. additional introducing the fine features of geometry of a CSU complex into the model, then the tightness of the connection in the first group is significantly increased. This suggests that systems with low fractal dimension, as more complex in structure, require multidimensional models of physical processes for their description.

In the quantitative ratio for the first group the viscosity growth rate of the oil is 1.6 times higher, and in absolute terms, with a fixed content of asphaltenes, the viscosity is 1.5-5.0 times higher than in the second group. So, experiments give an incompletely logical picture that with the same mass fraction of the dispersed phase, a higher internal resistance (shear stress) exists between the layers of the dispersion medium (liquid hydrocarbons) in the first group, although the shielding layer of the resin around the nucleus (see Fig. 4) here is more extended. That is explained by the low fractal dimension of the CSU nucleus.

Next, principal (theoretical) possibility of fractal geometry of aggregates effect on viscosity of the oil dispersed system is estimated. It was established by the numerical simulation method that hydrodynamic radius $R_h$ and radius of inertia $R_g$ of aggregates are related to each other by a simple relation

$$R_h = \beta_R R_g,$$  \hspace{1cm} (7)

where for three-dimensional relatively small spherical aggregates the value of $\beta_R$ is expressed in terms of fractal dimension $D$ by formula [34]

$$\beta_R = k^{1/2} \left[ \frac{2+D}{D} \right]^{1/2},$$ \hspace{1cm} (8)

where $k = 1.0…1.2$. It follows from (7) and (8) that for the first oil group the calculated hydrodynamic radius will be 1.61 times greater than the radius of inertia and 1.47 times for the second group.

The Einstein-Sima equation was used in order to evaluate the viscosity. The equation tells about cubic dependence of the viscosity $\eta$ from the hydrodynamic (Stokes) radius of particles:

$$\eta = \frac{4\pi N_a}{3M} R_h^3,$$ \hspace{1cm} (9)

where $N_a$ – Avogadro number; $M$ – molecular mass. In our case, increase in $R_h$ at the same radii $R_g$ aggregates in groups leads, according to (9), with constant other parameters, to the ratio of viscosities $\eta_1/\eta_2$ in groups equal to 1.3 units, which is close enough to the experimental estimate of 1.5 units obtained earlier for the lower limit of viscosity increase in the first group (see Fig. 6).

The simple ratio (8) no longer works in the region of the upper limit of viscosity, where the condition for the coherent state of asphaltene aggregates is satisfied for high-viscosity oil. It is theoretically impossible to estimate the ratio $\eta_1/\eta_2$.

Thus, decrease in fractal dimensions of the CSU nucleus leads to emergence of higher viscosity and, accordingly, rheological properties of oil at the same content of the dispersed phase in the form of asphaltenes. Hence, fractal asphaltene aggregates are more prone to association and formation of bulk supramolecular structures in oil.

**Temperature at which oil saturates with paraffin.** Temperature at which oil is saturated with paraffin (TOP) is the second technological parameter which characterizes phase transition of chain n-alkanes $C_{16}H_{34}$-$C_{36}H_{74}$ when the temperature decreases with the appearance of new
phase nuclei in the oil volume and formation of solid paraffin deposits. The higher the TOP, the greater the probability of premature formation of organic deposits of complex composition in production wells, sedimentation tanks, in-line and trunk pipelines, which leads to serious problems and accidents in extraction, transportation and refining of oil [35, 36].

As is known, saturation temperature under normal conditions depends on the content of paraffins, asphaltene and resins in oil. In this case, polar asphaltenes and resins can be simultaneously both propagators and inhibitors (terminators) of paraffin deposits [37, 38]. Experiments show, in that case TOP depends on the structure of CSU complexes.

Graphs in Fig. 7 shows dependence of the TOP on structural nanoparameter, which is equal to the ratio of thickness of the adsorption-solvate shell of resins to the radius of the core \( h_{ads}/r \), for two groups with a strong (graph 1) and weak (graph 2) asphaltene core. It can be seen from the figure that the dynamics of TOP differs significantly in selected oil groups.

It is found by the experiment that TOP varies according to linear equations if structural parameter CSU is varied

1st group – \( Y = 4.047X + 12.955, \rho = 0.909; \) \( (10) \)

2nd group – \( Y = 19.891X + 0.112, \rho = 0.829. \) \( (11) \)

According to (10) and (11), intensity of change in TOP in the second group is 4.9 times higher in comparison with the first one.

Thus, it can be said that TOP also depends on the fractal dimension of the asphaltene core in addition to all known parameters.

Correlation between the TOP and structural parameter \( h_{ads}/r \) is based on a mechanism that has several rules. Firstly, internal specific surface of adsorption of paraffin molecules increases with increase in size of the nucleus. The higher the number of adsorption centers the higher the probability of nucleation of critical embryos at a higher temperature and formation of a solid volumetric network of paraffin crystals. Secondly, when nuclear radius decreases the screening role of structureless resins increases, which act as blockers of inhibitory properties of asphaltenes with respect to paraffins.

Thus, for the second group with constant shell thickness (see Fig. 4) with changes in \( h_{ads}/r \) parameter from 0.5 to 1.3 units TOP value linearly increases to 31 °C (see Fig. 7) due to reduction in radius to 18.5 nm and decrease in temperature down to 10 °C in case of increase in size of nucleus in 2.6 times from the minimum to 48 nm, which corresponds to decrease in paraffin saturation temperature in 3.1 times.

Oil with a thicker nuclear solvation shell in the first group, by analogy, would seem to have a more tangible increase in growth of TOP due to screening compared to the second group. Nevertheless, experiments showed the reverse effect of decrease in both the growth rates and range of real values of TOP of oil.

The observed anomalous effect arises due to special properties of the low-dimensional fractal nucleus CSU, which manifest themselves in presence of long-range action (energy and power) asphaltenes on paraffin molecules. Asphaltenes of the core with shape of radial filamentary branches pierce the solvate shell of resins, are embedded in growing crystals of paraffin, which leads to a violation of their normal growth (size and shape) and further inhibits the process of approaching the crystals and their adhesion to form a solid solid paraffin wax, i.e. effect of inhibition.

Thus, low fractal dimensions of the nucleus stimulate the emergence of the inhibiting properties of asphaltenes with respect to paraffin hydrocarbons. High fractal dimensions reduces and blocks them.
In practice, the established linear dependence of TOP on structural parameters of CSU (see Fig. 7) opens the possibility of direct control of the process of precipitation of paraffin hydrocarbons during the cooling of oil in tubing or pipeline through the formation of the necessary stable structure of asphalt-shadow complexes due to both a decrease in the thickness of the solvate shell and increase in the radius of the asphaltene core.

**Temperature of aggregation of asphaltenes.** Since spontaneous increase in sizes of active asphaltene complexes is one of the main reasons for formation of ARP in various technological processes influence of temperature on processes of aggregation of asphaltenes in native oil is considered [39].

As is known, increase in temperature leads to increase in velocity of Brownian motion of molecules and disintegration of bound associative complexes, flocculae, aggregates and chemical bonds in liquid and solid substances, i.e. to destruction of existing order in matter.

During extraction, transportation and storage of petroleum products, technological processes associated not with growth but with slow decrease in temperature of the liquid in a tubing or pipeline. This finally leads to decrease in the aggregative stability factor of the oil dispersed system, which also depends on the fractal dimension of the CSU complexes.

Laboratory temperature studies of native oil were conducted using a technology close to the commercial. Initially, the oil sample was heated above the TOP to 60 °C and held for 4-5 hours with periodic mixing in a thermostat until the asphaltene complexes were completely destroyed and dissolved in oil. Then the sample was placed in a sealed thermostat container, temperature of the oil in it was gradually decreased in steps of 5 degrees in the range from 60 to 3-5 °C and held at each stage for 40 minutes. During the such slow step-by-step lowering of the temperature by NMR, restructuring of the nanostructure of the CSU of native oil due to change in mean radius of the asphaltene core was monitored.

The Fig. 8 shows an example of temperature dependences of the growth dynamics of the CSU nucleus radius in aggregation of asphaltene particles for two viscous samples of native oil. It can be seen that in the entire range from 60 to 5 °C a close to linear inverse model of aggregation is performed, which is characterized by an average angular coefficient (aggregation coefficient) \( \alpha = dR/dT \) \( (R – radius of the nucleus, T – temperature) \) of the evaluation line. This parameter quantitatively reflects the intensity of the effect of uniform oil cooling on the resulting structural size of the CSU nucleus during the aggregation of asphaltene particles.

![Fig. 8. Dynamics of change in radius of asphaltene core from temperature and angle of inclination for viscous samples of native oil of Mosinskoe (1) and Kanakhinskoe (2) fields](image)

The experiments conducted showed that aggregation processes have characteristic features that make up from the fractal dimension of the nucleus of native oil. In particular, decrease in fractal dimension leads to an increase in aggregation rate of asphaltene particles, i.e. increase in angular coefficient of the kernel growth chart. Thus, as it seen from the Fig. 8, aggregation coefficient for the Well 235 of the Mosinskoe oil field from the first group with lower \( (D = 1.556) \) nucleus dimension is equal to 0.499 nm/degree, and 0.297 nm/deg for the Well 71 of Kanakhinskoe oil field of the second group \( (D = 2.040) \). In general, average aggregation coefficient \( \alpha_{av} \) for the first group of native oil is 3.6 times higher than the coefficient \( \alpha_{2av} \) in comparison with the second group \( (0.391 against 0.110 nm/degree) \).

The features obtained are explained by the fact that low-dimensional structure of CSU nucleus has internal adsorption surface and surface energy controlling its reactive and solvating activity much higher than in high-dimensional nucleus due to...
more branched, flexible and mobile asphaltene framework. That framework is capable of conformational transformations, which reduces steric hindrances and enthalpy of association upon contact and “clumping” of aggregates such as “particle-cluster” or “cluster-cluster” and causes real increase in intensity of aggregation with decrease in oil temperature.

An additional regularity has been established. The process of aggregation of asphaltene particles depends both on fractal dimensions and initial size of the nucleus-beam (growth center) from which aggregation begins. In case the equilibrium radius of the nucleus is taken as the starting point for oil samples at room temperature (25 °C) then the picture below is experimentally obtained.

For oil of the second high-dimensional group, as the radius of the initial growth center increases, the coefficient of thermal aggregation $\alpha_2$ (Fig. 9, graph 2) decreases monotonically, which is well described by an equation of the form

$$Y = -0.1732\ln X + 0.7902, \rho = 0.952. \ (12)$$

That means the aggregation of asphaltenes into relatively small clusters occurs with a higher intensity ($\alpha_2 = 0.37 \text{ nm/deg at } R = 10 \text{ nm}$) than for larger ones ($\alpha_2 = 0.10 \text{ nm/deg at } R = 55 \text{ nm}$).

Having temperature drop in the range from 55 to 5 °C, the average absolute increment of $\Delta R$ in large aggregates is 5.5 nm and 20.4 nm in small ones. In other words.

Large aggregates have a relatively small increase in radius by 15-18 % while the size of the nucleus is significant and amounts to 150-200 %, i.e. significantly exceeds the initial seed size.

The monotonous decrease found in aggregation $\alpha_2$ is explained by the fact that at the equilibrium radius of the nucleus from 10 to 45 nm, firstly, linear growth of the potential barrier is observed – activation energy from 7.5 to 17 kJ/mol (see Fig. 3, graph 2), which inhibits and limits the aggregation of particles. Secondly, experiments show that the solvate number $Z$ decreases from 1.6 to 1.05 units as the diameter of the adsorption center increases, i.e. the number of contacting particles in the first coordinate sphere of the nucleus. Superposition of that negative parameters of the kinetic factor leads to a decrease in the probability of approach and subsequent collision of particles with the nucleus, which is responsible for the expected decrease in the aggregation coefficient with increase in the equilibrium radius.

Thus, in spite of the small content of asphaltenes in 0.3-1.3 % by weight in a light-weight oil of the second group, precipitation of ARP due to the increased temperature aggregation will be formed in the tubing quite intensively, which is confirmed by numerous commercial data. At the same time, phase transition of paraffins is the main reason for precipitation of ARP in heavy and viscous oil. Thermal aggregation of asphaltenes is shifted to the background due to extremely low contribution.

The monotonic character of the dependence established by (12) (see Fig. 9, graph 2) confirms the presence in the second group of native oil of a more homogeneous porosity nucleus, which becomes similar (in terms of its properties) to bodies with a constant density.

Considering the process of aggregation for the first group of oil, as was shown above (see Fig. 3, 4), the oil of a low-dimensional group is characterized by the emergence of structural and energy extremal states. This feature was also established by us during the study of thermal aggregation of asphaltenes. In particular, with initial increase in beam radius in the light oil of the first group, intensity of aggregation of asphaltenes does not decrease as in the second group, on the contrary, increases sharply, reaching a maximum of 0.5 nm/deg at the maximum
Analysis of experimental data shows that anomalous (reverse) in direction aggregation at the initial seed interval from 12 to 18 nm is realized when the synergistic action of the the components of the kinetic factor is combined such as presence of a local minimum (15 kJ/mol vs. an average of 19 kJ/mol) of activation on the border with asphaltenes (see Fig. 3, graph 1), local minimum (15.5 nm vs. average at 26.6 nm) thickness of the solvate shell (see Fig. 4, graph 1) and high (Z = 1.7 units) of the solvate number of the nucleus. Each of them determines the probability of implementing a separate stage of a complex dynamic aggregation process, and the product of the probabilities of physical quantities determines the final probability of the process as a whole. Thus, a solvate number growths and provides a statistically significant number of active particles near the nucleus, a low activation barrier facilitates the approximation of aggregating particles and small thickness of the protective shell facilitates direct contact of the particle with the nucleus and formation of a new stable structure of the aggregate. Eventually, kinetic energy of the reacting particles is converted into the potential energy of interparticle bonds, and temperature aggregation index increases by an abrupt increase of 25 %, although, it should decrease slowly with increasing beam sizes. It should be noted that the process observed characterizes only fractal asphaltenene aggregates with a low core dimension.

Thus, as the fractal dimension of the nucleus decreases, the number of possible microstates (degrees of freedom) of the system increases, which, according to L.E. Boltzmann corresponds to increase in system's entropy. Increase in entropy reflects the fact that the structure of CSU became less ordered (irregularity, chaos), which plays a role of generator of the high extremal activity of the CSU nucleus, which controls the macroproperties of native oil, decreases in order.

Quantitative estimates carried out from the data in Fig. 9 show that for the oil of the first group in subcritical mode, the slope of the angle $d\alpha_i/dR$ of the left branch of the graph 1 describing the anomalous aggregation is $+16.7 \cdot 10^{-3}$ fractions of unit, and in the overcritical region of the normal decay, respectively, $13.3 \cdot 10^{-3}$ fractions of units. At the same time for the second oil group, the
analogous decline \( \frac{d\alpha_2}{dR} \) of aggregation of asphaltenes is \(-6.25 \cdot 10^{-3}\) fractions of unit, which is 2.1 times less than for the first group.

It follows that oil with a lower fractal dimension of the nucleus is more prone to aggregate and form the ARP deposits when temperature is decreased.

**Effectiveness of ARP deposit inhibitors**

Let's have a look at the results of experiments on the direct observation of ARP deposits on a metallic surface. One of the methods for preventing deposition of AFS is chemical inhibitors.

When an inhibitor, which is a surfactant, is dissolved in oil, adsorption occurs on the interfacial surface of the CSU, which leads to a change in the internal structure of the liquid and transition of oil dispersion system into a new thermodynamic state. In particular, we showed in [11] that an effective inhibitor leads to a decrease (peptization) of the mean radius of the asphaltene CSU nucleus and increase in activation energy \( E_a \) of molecules of dispersion medium. In order to estimate the weight of ARP deposits on the metal surface after oil treatment with inhibitors, we obtained an analytical equation [40] that relates the mass of paraffin deposits to energy parameter \( \Delta E_a \) of liquid molecules:

\[
Y = 1 - U\Delta E_a, \tag{14}
\]

where \( Y \) – relative mass equal to the ratio of weight of ARP matter after treatment to mass of deposits before treatment with oil inhibitor, \( Y = m_{\text{inh}}/m_{\text{ini}} \), \( U \) – oil constant; \( \Delta E_a \) – difference \((E_{a\text{inh}} - E_{a\text{ini}})\) activation energy with and without an inhibitor.

An efficiency coefficient \( E \) used in practice is calculated using the equation (14), that shows relative decrease in weight of ARP deposits on the metal surface after oil is being treated with an inhibitor:

\[
E = 1 - Y = U\Delta E_a. \tag{15}
\]

Thus, the effectiveness of the inhibitor is proportional to the difference in the activation energy of \( \Delta E_a \) hydrocarbons with a positive sign, showing how much this inhibitor can increase the potential barrier for dispersion medium molecules that directly controls the growth and adhesion of paraffin crystals.

The Fig. 10 shows the results of the cold plate method of laboratory experiments testing five different inhibitors of ARP deposits of SNPCH using samples of two oil groups differing in fractal dimensions of the asphaltene nucleus.

Firstly, it can be seen that the experiments confirm the theory (15) on example of use of different in physical and chemical properties of oil and chemical reagents. The obtained dependences on the effectiveness of the ARP deposits inhibitors are described by linear equations which are as follows:

1\textsuperscript{st} group

\[
E = -0.0755\Delta E_a + 0.015, \quad \rho = 0.992, \tag{16}
\]

2\textsuperscript{nd} group

\[
E = -0.0480\Delta E_a + 0.021, \quad \rho = 0.999. \tag{17}
\]

Secondly, a clear difference in dynamics of relative weight of ARP deposits by oil groups was obtained. In particular, the intensity of \( dE/d\Delta E_a \) of inhibition of ARP deposits in the first group is 1.6 times larger than in second one, which exactly coincides with the analogous coefficient when comparing the dynamic viscosity (see Fig. 6).

Third, oil with low fractal dimensions shows a higher positive efficiency when treated with different inhibitors. So, for example, comparing the inhibitor that increases activation energy by +4 kJ/mol, the average efficiency of reducing the
weight of the ARP deposits for the second oil group is 21.3 %, and 31.7 % in the first one which gives a difference of 48.8 % rel.

According to the negative effect (see Fig. 10), which often leads to increase in weight of ARP deposits when treated with inhibitors, as compared to a blank sample, the first group with a low-dimensional CSU nucleus is also leading.

A negative effect, especially in the case of heavy viscous oil, arises when the chemicals start in the disperse system the negative growth processes of the radius of the asphaltene CSU nucleus and reduce threshold energy of oil activation, which gives a negative sign of difference in $\Delta E_a$ in (15), which is generally stimulating more intensive growth of paraffin crystals and increase in their mass with adhesion on the surface of equipment [6].

As a result of the set of studies carried out, it was shown that the change in the fractal dimension of asphaltene aggregates of CSU due to external factors and physical and chemical fields can cause abnormal behavior of oil's technological properties (viscosity, mobility, TOP, aggregation of asphaltenes and precipitation of ARP deposits), which is not a subject to the additivity law. The established correlation of nano-properties and macro-characteristics of native oil in terms of methodology allows more reasonably searching for and developing new, including non-traditional, methods for increasing the efficiency of technological processes in oil and gas industry.

The results obtained in the work can be used in practice while improving modern methods of controlling the properties of oil dispersed systems in development, production and refinery of hydrocarbons.

Conclusions

1. A complex analysis of energy and structural features of the structure of asphaltene nanoaggregates was carried out experimentally and fractal dimensions of CSU nucleus were calculated for two alternative groups of native oil, which is 2.040 and 1.556 units.

2. It is established that oil with low fractal dimensions of the nucleus has a higher (by 1.5-5.0 times) dynamic viscosity, and theoretical substantiation of this physical phenomenon is carried out.

3. It is revealed that oil with low fractal dimension due to its structural features has an increased natural inhibition ability of the asphaltenes of the core in relation to growth and precipitation of solid paraffins forming ARP deposits.

4. According to the data of temperature studies, it is established that dynamics of growth of the radius of the CSU nucleus during the aggregation of asphaltene particles is well described by the linear model, while the aggregation intensity is on the average higher by 3.6 times in oil with a lower fractal dimension.

5. It is additionally determined that the coefficient of thermal aggregation of asphaltenes depends on the radius of the initial seed and for oil with a high fractal dimension is described by a monotonic logarithmic model.

6. Anomalous nonlinear aggregation is established for oil with low dimensions at small radii of the initial beam, which violates the monotonous nature of change in aggregation coefficient.

7. According to results of the experiments, it was established that effectiveness of action of the ARP inhibitors in oil with low fractal dimension is 5-49 % higher compared to the group of high-dimensional oil.

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