INTRODUCTION

It is known that when developing the methods for microbial utilization of various pollutants, including the components that make up drilling fluids and cuttings, it is advisable to screen for degrading microorganisms in contaminated habitats, because under such conditions, under the influence of selective pressure, microbial strains with a high destructive potential are formed.

The presence of microorganisms in oil reservoirs has long been established. On the example of various oil-bearing horizons, the existence of various metabolic groups of anaerobic microorganisms in the layers was established, i.e. the sulfate-reducing bacteria that reduce thiosulfate, Fe^{3+} and elemental sulfur, fermentative bacteria, acetogens and methanogens. Oil-bearing horizons, as a rule, are characterized by anoxic conditions, thus attracting the main attention of researchers. However, aerobic microorganisms also live in oil reservoirs, where they enter with injected water, drilling fluid during field development and in connection with the natural hydrodynamics of groundwater. In exploited oil fields, natural hydrocarbon-oxidizing microorganisms (hereinafter referred to as HOM) are the initial link in the microbial trophic chain that biodegrades oil.

The exploitation of oil fields, which until the moment of their development were practically not exposed to external influences, is accompanied by the entry of surface microorganisms, oxygen and some biogenic elements into the reservoir, which creates prerequisites for the development of specialized microbial communities, one of the main components of which are HOM.

MATERIAL AND METHODS

At present, a wide distribution of microorganisms in the waters of oil fields has been shown. Many authors searched for viable natural
hydrocarbon-oxidizing microorganisms in the groundwater of deep wells (1900 and 3200 m), which were calcium chloride thermal brines.

In this biotope, which is specific in terms of environmental factors, it was possible to find UOM and fungi. Drill cuttings (hereinafter referred to as BS) that have been at specialized sites of oil fields for a long time, can also serve as selective agents for the formation of HOM, as well as bacteria-destructors of various components of drilling fluids (Benka-Coker, Olumagin, 1995).

In connection with the foregoing, the microbial communities of several BS samples (BS 1 and BS 2) with a high pH value (pH 9) and a degree of mineralization (15%) were studied to identify microorganism-destructors of chemical reagents that are part of drilling fluids.

In the course of the studies, it was shown that the number of heterotrophic microorganisms isolated by direct seeding in the microbial communities of the analyzed samples of BS 1 and BS 2 was 1×10^3 and 8.9×10^3 cells/g, respectively (Table 1). Only aerobic bacteria of microbial communities were studied, because it is preferable to use aerobic bacteria for microbial utilization of BS located in special areas (sites, barns, etc.).

In connection with the multicomponent composition of BS, which includes, in addition to the hydrocarbon base, various chemical compounds, such as surfactants (hereinafter referred to as surfactants), starch, carboxymethyl cellulose (hereinafter referred to as CMC), etc., for a more detailed characterization of microbial communities, BS was evaluated in terms of the number of bacteria that hydrolyze starch, CMC, as well as microorganisms-destructors of nonionic surfactants (hereinafter referred to as nonionic surfactants), anionic surfactants (hereinafter referred to as surfactants) and natural hydrocarbon-oxidizing microorganisms (hereinafter referred to as HOM).

The content of microorganisms of various physiological groups in drill cuttings are presented in Table 1.

As it was established (Table 1), the BS 2 sample contained: UOM – 6.1×10^3 and 8.8×10^2 cells/g of microorganisms that hydrolyze starch. At the same time, microorganisms-destructors of sintanol, sulfonol and CMC were not detected. The microbial community of sample BSh 1 did not differ in diversity; here, HOM, destructors of nonionic surfactants, surfactants, CMC and starch were not detected in a noticeable amount.

When bacteria were isolated by using the method of enrichment cultivation, hydrocarbon substrates were used as the only source of carbon and energy, to which, presumably, the BS microorganisms could be partially adapted; these are drilling fluids based on mineral oil (BR 1) and diesel fuel (hereinafter – DF) (BR 2). The data on the total number of heterotrophic microorganisms and the content of HOM, microorganisms-destructors of nonionic surfactants, surfactants, as well as the bacteria that hydrolyze starch and CMC, obtained from the analysis of enrichment cultures using the BS samples, are presented in Table 1.

The results of the experiments showed that the addition of selective food sources to the cultivation medium increased the number of bacteria

| Samples | Extraction methods | Heterotrophic | UOM | Hydrolyzing starch | Hydrolyzing CMC | Destructors nonionic surfactants | Destructors AS |
|---------|-------------------|---------------|-----|-------------------|----------------|---------------------------------|----------------|
| BSh 1   | 1. Direct seeding  |               |     |                   |                |                                 |                |
|         |                   | 1.0×10^3 ±0.30| <50±0.44 | <50±0.00          | <50±0.00       | <50±0.00                        | <50±0.00       |
|         | 2. Cumulative      | 2.8×10^5 ±0.58| 4.4×10^4 ±0.22 | <50±0.00       | <50±0.00       | <50±0.00                        | <50±0.00       |
|         | cultivation (+BR 1) |                   |     |                   |                |                                 |                |
|         | 3. Cumulative      | 2.0×10^4 ±0.22| 1.1×10^3 ±0.44 | 4.3×10^2 ±0.42 | <50±0.44      | <50±0.00                        | <50±0.00       |
|         | cultivation (+BR 2) |                   |     |                   |                |                                 |                |
| BSh 2   | 1. Direct seeding  |               |     |                   |                |                                 |                |
|         |                   | 6.9×10^3 ±0.38| 6.1×10^2 ±0.34 | 8.8×10^2 ±0.46 | <50±0.54      | <50±0.00                        | <50±0.00       |
|         | 2. Cumulative      | 1.4×10^3 ±0.38| 3.3×10^2 ±0.45 | 6.9×10^2 ±0.58 | <50±0.00      | <50±0.00                        | <50±0.00       |
|         | cultivation (+BR 1) |                   |     |                   |                |                                 |                |
|         | 3. Cumulative      | 1.8×10^3 ±0.40| 1.6×10^2 ±0.28 | 4.2×10^2 ±0.52 | <50±0.00      | <50±0.00                        | <50±0.00       |
|         | cultivation (+BR 2) |                   |     |                   |                |                                 |                |

Note: “<50” means that when bacteria were seeded from a 1:5 dilution, microorganisms of the corresponding groups were not detected.
of certain above-mentioned groups compared with the direct seeding of microorganisms from BS. In the mud sample BSh 1, the total content of heterotrophic bacteria increased 10 times when it was cultivated with drilling fluid based on diesel fuel and significantly higher – by two orders of magnitude – with a solution based on mineral oil. The use of these oil-based drilling fluids (hereinafter referred to as OBM) as nutrient substrates contributed, to a large extent, to the development of the UOM group. With direct seeding from this sample, HOM could not be detected, whereas, with accumulation cultivation, the number of HOM reached the following values: 1.1×10⁴ and 4.4×10⁴ cells/ml when drilling fluids based on diesel fuel and mineral oil were introduced into the medium, respectively. In the BSh 1 sample, compared with direct seeding, the number of microorganisms that hydrolyze starch, when cultivated with a model solution based on diesel fuel (BR 2), increased by 2 orders of magnitude, while the addition of drilling fluid based on mineral oil (BR 1) did not affect this group of bacteria.

In the BSh 2 sample, the total number of heterotrophic microorganisms, as well as in BSh 1, reached 10⁵ cells/ml when it was cultivated with solutions of BR 1 and BR 2, the number of UOM was 10⁴ cells/ml, the number of bacteria that hydrolyze starch. Both drilling fluids, which we used as selective reagents, equally contributed to the development of microorganisms of different groups contained in the BSh 2 sludge, in contrast to the BSh 1 sample, where the drilling fluid based on mineral oil stimulated the growth of heterotrophic bacteria and UOM to a greater extent.

It should be noted that the additions to the cultivation medium in the form of drilling solutions did not stimulate the development of bacteria-destructors of nonionic surfactants, surfactants, and CMC in both BS samples, which may be due to a rather low concentration of these reagents in the solutions or their toxicity.

Thus, it can be noted that aerobic heterotrophic bacteria are contained in BS in different amounts, which is probably due to the specific environmental conditions, as well as the different composition of the cuttings and, as a result, the presence of various sources of nutrients. Among the bacteria included in the BS communities, UOM, starch hydrolyzing microorganisms, are found. This creates the prerequisites for the isolation of BS reagents and drilling fluids from such communities of active destructors, including those hazardous to the environment.

RESULT

Morphological and physiological-biochemical characteristics of isolated cultures

As a result of the experiments, 7 pure microbial cultures were isolated from BS, which were later studied in more detail.

By direct seeding, a strain with the laboratory code NSh was isolated from the BSh 1 sample, and OBR 1 was isolated from the BSh 2 sample.

Figure 1. Images of cells of isolated cultures at the age of 24 hours, obtained using transmission electron microscopy
Using the enrichment cultivation method, from the BSh 1 variant, where the drilling fluid BR 2 was the only source of carbon, samples were isolated (hereinafter referred to as the OBR): OBR 3.1, OBR 3.2, and OBR 3.3; when using drilling fluid BR 1, OBR 1.1 was isolated from this sample; from sample BSh 2 with drilling fluid BR 2 – OBR 4.1.

In order to identify the isolated strains, research was carried out to study their cultural-morphological and physiological-biochemical characteristics. The results of the analyses performed are presented in tables 2 and 3.

The main diagnostic morphological and morphometric characteristics of microbial cells were obtained using electron microscopy (Figure 1).

On the basis of the studies carried out, the cultures isolated from drill cuttings were identified based on the results of a comparative analysis of cultural-morphological, morphometric and physiological-biochemical characteristics in accordance with the criteria for bacterial differentiation proposed in the 9th edition of the manual “Burges Key to Bacteria” (1997), as well as in accordance with the principles molecular typing of prokaryotic cells as: Bacillus circulans NSh; B. firmus OBR 1.1; B. firmus OBR 3.1; Solibacillus silvestris 3.2, B. circulans OBR 3.3, Halomonas sp. ABR 1 and Erwinia rhapontici ABR 4.1.

Therefore, from the BS 1 sample by direct seeding and with the help of accumulative cultivation, the following were isolated: Bacillus

| Laboratory code of cultures | Microscopic study | Growth on MPB | Growth on MPA | Relation to oxygen | Temperature optimum | Characteristics of the colonies |
|-----------------------------|-------------------|--------------|--------------|--------------------|---------------------|-------------------------------|
| NS                          | Gram-variable rods: at the age of 24 hours - gram-negative long or slightly curved, located singly and in pairs; 48 h - gram-positive short. Endospores are ellipsoidal, lying subterminally. Spores are cylindrical. | When incubated for 24-48 hours without shaking there is a uniform turbidity of the broth, the formation of a film, sediment | Good growth on the first day at 28°C | Facultative anaerobe | 20-30°C | Opaque cream with a slightly convex shiny surface and smooth edges; d colonies - 1.0-3.0 mm |
| OBR 1.1                     | Gram-positive rods, located singly and in the form of chains of two or three cells. Endospores are ellipsoidal, lying subterminally. Controversy. Situation proposed in the 9th edition of the manual | When incubated for 24-48 hours without shaking, a precipitate is observed with a uniform clouding of the environment | Good growth on the first day at 28°C | Facultative anaerobe | 20-30°C | Opaque cream with a slightly domed shiny surface and smooth edges; d colonies - 2.0-4.0 mm |
| OBR 3.1                     | Gram-positive rods, located singly. Dispute does not form | Precipitation occurs when incubated for 24-48 hours without shaking | Good growth on the first day at 28°C | Facultative anaerobe | 20-30°C | Opaque white with a slightly convex shiny surface and smooth edges; d colonies - 2.0-3.5 mm |
| OBR 1                       | Gram-negative straight rods located singly. Dispute does not form | When incubated for 24-48 hours without shaking, film formation is observed on the surface | Good growth on the first day at 28°C | Aerobe | 20-42°C | Round shape, with a rough surface, shiny, white-cream color with smooth edges, dry consistency; d colonies - 2.0-4.0 mm |
| OBR 3.2                     | Gram-positive straight rods located singly. Generates controversy. Endospores are spherical, lying terminally in swollen sporangia | When incubated for 24-48 hours without shaking, film formation is observed on the surface of the medium | Good growth on the first day at 28°C | Aerobe | 20-35°C | Small rounded, smooth, shiny, white, homogenous, with smooth edges, mucous consistency; d colonies - 1.0-3.0 mm |
| OBR 3.3                     | Gram-positive long rods, located both singly and in pairs. Generates controversy. Endospores are ellipsoidal, lying subterminally. Controversy cylindrical. | When incubated for 24-48 hours without shaking, a precipitate is observed with a uniform turbidity of the environment | Good growth on the first day at 28°C | Facultative anaerobe | 20-30°C | Opaque white with a smooth luster surface and smooth edges; d colonies - 1.0-3.0 mm |
| OBR 4.1                     | Gram-negative straight rods with rounded ends, located singly. Does not form a dispute | When incubated for 24-48 hours without shaking precipitate formation is observed | Good growth on the first day at 28°C | Facultative anaerobe | 20-42°C | Large rounded, smooth, shiny, white, opaque, homogenous, with smooth edges, mucous consistency; d colonies - 3.0-5.0 mm. After 48 hours of incubation, the colonies become slightly pink shade. |

Table 2. Results of the performed analyzes
The following cultures were isolated from the BSh 2 sample: *Halomonas* sp. ABR 1 and *Erwinia rhapontici* ABR 4.1.

So, from the sample BS 1 by direct seeding and with the help of accumulative cultivation, the following were isolated: *Bacillus circulans* NS; *B. firmus* OBR 1.1; *B. firmus* OBR 3.1; *Solibacillus silvestris* 3.2 and *B. circulans* OBR 3.3. The following cultures were isolated from sample BSh 2: *Halomonas* sp. ABR 1 and *Erwinia rhapontici* ABR 4.1.

Saprotrophic spore-forming bacteria (bacilli) are common inhabitants of soils; they are actively involved in the processes associated with the decomposition of organic substrates. It has been shown that under the conditions of oil pollution in soils, active reproduction of bacilli cells occurs, and species characteristic of soils with weak mineralization processes multiply to a large extent, and the diversity of the species composition of the spore-forming microbiota increases (Kireeva and Rafikova, 2007).

The authors (Bento et al., 2003) isolated from the soils contaminated with diesel fuel (California, USA) a HOM consortium based on *Bacillus cereus*, *Bacillus sphaericus*, *Bacillus fusiformis*, *Bacillus pumilus*, *Acinetobacter*.

Table 3. Physiological and biochemical properties of isolated cultures

| Signs                  | Laboratory code of cultures |
|------------------------|-----------------------------|
|                        | NS  | OBR 1.1 | OBR 3.1 | OBR 3.2 | OBR 3.3 | OBR 1  | OBR 4.1 |
| Acid resistance        | -   | -       | +       | +       | -       | -      | -       |
| Mobility               | +   | +       | +       | +       | +       | +      | +       |
| Oxidases               | -   | -       | -       | -       | -       | +      | -       |
| Catalase               | +   | +       | +       | +       | +       | +      | +       |
| H₂S                   | +   | +       | +       | +       | +       | +      | +       |
| Indole                 | -   | -       | -       | -       | -       | -      | -       |
| NH₃                   | -   | -       | -       | +/-     | -       | +/-    | +       |
| Nitrate reductase      | +   | +       | +       | -       | -       | -      | +       |
| Phenylalanine deaminase| +   | -       | -       | -       | -       | -      | -       |
| Casein                 | +/- | -       | +/-     | +/-     | +/-     | +/-    | -       |
| Gelatinase             | +/- | -       | +       | -       | +/-     | -      | +/-     |
| Arginine hydrolases    | -   | -       | -       | -       | -       | +/-    | -       |
| Lysine decarboxylases  | -   | -       | -       | -       | -       | +/-    | -       |
| Ornithine decarboxylases| -  | -       | -       | -       | -       | -      | -       |
| Lecithinase            | -   | -       | -       | -       | -       | -      | -       |
| Ureas                  | -   | -       | -       | -       | -       | -      | -       |
| Hydrolysis of starch   | +/- | +       | -       | +/-     | -       | +/-    | -       |
| Utilization of citrate | -   | -       | -       | +/-     | -       | -      | -       |
| Growth at 10 °C        | -   | +       | -       | -       | -       | -      | -       |
| Growth at 42 °C        | +   | +       | +       | +       | +       | +      | +       |

| Oxidizes               |                  |
| Glucose                | OF | OF | OF | O | OF | O | F |
| Lactose                | F  | OF | F  | O | OF | O | - |
| Arabinose              | OF | O  | -  | O | OF | - | F |
| Sucrose                | -  | O  | OF | O | OF | O | O |
| Maltose                | -  | O  | O  | O | O  | O | - |
| Mannitol               | O  | O  | OF | O | F  | O | - |
| Xylose                 | O  | O  | -  | O | O  | - | - |
| Fructose               | O  | -  | -  | - | O  | - | - |
| Galactose              | O  | O  | -  | O | O  | O | F |
| Voges-Proskauer reaction| - | -  | -  | - | -  | - | - |

Note: “O” – oxidation, “F” – fermentation, “-” – no reaction.
It is known (Ne-trusov and Kotova, 2006; Gosmanov et al., 2011) that the formation of spores for bacteria is a factor in the preservation of the species under unfavorable conditions (changes in temperature, pH of the environment, lack of nutrients, the presence of toxins, etc.).

The fact that the representatives of spore-forming microorganisms dominated among the bacteria isolated from BS indicates their wide adaptive potential, which is associated, among other things, with spore formation.

Substrate spectrum of bacteria isolated from drill cuttings

Due to the fact that the hydrocarbon component of OBM poses a great environmental hazard to the environment as well as requires rapid and complete destruction, the bacteria isolated from the microbial communities of BS were studied for the ability to degrade petroleum hydrocarbons.

When studying the destructive potential of microorganisms isolated from BS, as the only source of carbon and energy, crude oil, vaseline oil, diesel fuel, drilling fluid based on mineral oil, a number of individual hydrocarbons were added to the agar mineral medium: n-alkanes (hexane, heptane, octane, decane, hexadecane), aromatics (benzene, toluene, xylene, cumene, nitrobenzene), alicyclic compounds (cyclohexane) and PAHs (naphthalene).

All studied bacteria were characterized by a distinct growth on all used oil products (Figure 2), with the exception of the S. silvestris OBR 3.2 strain, whose growth was not observed in the variants with vaseline oil and diesel fuel.

Growth on oil and oil products suggests that bacteria have appropriate enzyme systems for the degradation of hydrocarbons and mechanisms to suppress the toxic effect of oil. Oil contains thousands of substances in its composition, and the share of a few of them reaches at least one percent. Obviously, for effective oil degradation, the strain must be able to compensate for the toxic effect of the pollutant and be able to use at least some of the oil components as sources of carbon and energy.

On individual hydrocarbons, the studied microorganisms grew with different intensity. Among them, several crops can be distinguished (B. firmus OBR 1.1; B. firmus OBR 3.1; B. circulans OBR 3.3 and B. circulans NS), which actively used all the oil products and individual hydrocarbons were tested for growth (Figure 3). The relatively greater width of the spectrum of hydrocarbons consumed by these microorganisms can be interpreted as their relatively greater ability to consume pollutant oil products.

All the studied bacteria were unable to assimilate cyclohexane as the only source of carbon, which indicates the high toxicity of naphthenic hydrocarbons for these microorganisms. Bacteria also did not grow on a mineral medium with naphthalene, which acts as the only source of carbon and energy, at the same time, on MPA with naphthalene at a concentration of 0.4 mg/l, all cultures showed good growth on the first day, which indicates the stability of the data bacteria to naphthalene at this concentration.

Melnikov and Karaseva (2005) reported that the bacterial strains isolated from oil-contaminated soil most often grew on n-alkanes.
(14.5–33.3%), much less frequently on aromatic hydrocarbons (0–4.3%).

It should be noted that, despite the high toxicity of aromatic hydrocarbons, in particular, nitrobenzene (Taran, 2012; Elder and Kelly, 1994), the microbial strain *B. firmus* OBR 1.1 was characterized by a barely pronounced growth on nitrobenzene and distinct growth on toluene. The *B. firmus* OBR 3.1, *S. silvestris* OBR 3.2, and *B. circulans* OBR 3.3 microorganisms were characterized by distinct growth on the medium with benzene.

Thus, the obtained results testified to the diversity of trophic possibilities of the isolated microorganisms in relation to oil products and individual hydrocarbons, components of BS.

**Quantitative assessment of the destruction of petroleum products by the bacteria isolated from drill cuttings**

It is known that the most common dispersion media in OBM are oil, diesel fuel and mineral oil (Soloviev, 2003; Smirnova, 2011; Young, 1994). Therefore, the destructive activity of the isolated microorganisms was studied in relation to these dispersion media (petroleum products), which were added to the M9 mineral medium as the only source of carbon and energy at a concentration of 0.4% (by weight).

The destructive activity of bacteria was determined after 10 days of cultivation, according to the residual content of petroleum hydrocarbons in the mineral medium by using the gravimetric method (RD 52.18.647-2003. Methodological instructions..., 2003).

After 10 days, in the control samples (without the addition of bacteria), a slight decrease in the studied dispersion media was observed, which is due to physicochemical processes occurring over time in the medium with oil products (photolysis, hydrolysis, etc. without the participation of biota) (Ivanenko, 2006; Shukla, 1990). The amount of oil at the same time decreased by 7, mineral oil – by 8, diesel fuel – by 15%.

The destructive activity of bacteria was 28–40% in different strains. The maximum destructive activity towards oil was observed in strains *B. firmus* OBR 3.1 (Figure 4a) and *B. circulans* NSh (38%) (Figure 4b), as well as in *Halomonas* sp. OBR 1–40%.

Differences are visualized in Figure 4. In a liquid medium with oil as the only source of carbon and energy and microbial strains, there is no oiliness on the walls of the flask, the medium becomes cloudy due to bacterial growth; the oil is in a dispersed state, in the form of a finely dispersed emulsion, in granules of different sizes. In the control flask without bacteria, the medium is transparent; the oil is without any changes in color and state of aggregation. The fact of oil dispersion suggests that these strains under certain cultivation conditions are able to synthesize and secrete surfactants that emulsify oil.

If the destructive activity of bacteria isolated from BS is compared with the activity of other oil degrading cultures, the following can be noted: in a model experiment Loginova et al. (2004) showed that the biological product “Bacispecin” based on the strain *Bacillus* sp. 739 is able to utilize oil in 40–60% in 30 days, depending on the degree of pollution. In the work of Klyuyanova (2009) it was found that the *B. firmus* SDS-1 strain isolated...
from oil-contaminated soil is able to utilize oil in a liquid medium in up to 82% in 10 days of cultivation. Plotnikova (2010) showed in her studies that *Halomonas* sp. is able to assimilate aromatic and polyaromatic hydrocarbons, which are contained in the composition of oil, up to 55%. The bacterial strains isolated by Ilyina et al. (2003) from oil-contaminated soils showed from 21% to 30.5% destruction of total petroleum hydrocarbons in 15 days of cultivation. They were selected as the best among the other 30 isolates. The biological product “COBE-10” was created based on them. Among these strains were: *Bacillus* sp., *Rhodococcus* sp., *Providencia* sp., *Citrobacter* sp.

Thus, the destructive activity of the isolated microorganisms in relation to petroleum hydrocarbons, although not too high, is at the same time comparable with the literature data.

In relation to DT, two strains, i.e. *B. circulans* OBR 3.3 and *B. circulans* NSh, showed a noticeable destructive activity. The total content of hydrocarbons during the cultivation of these strains decreased by 35.5 and 25%, respectively. The results obtained are consistent with the literature data. In the experiments of Astrov et al. (1998) showed a high destructive activity of the *Bacillus* sp. strains, which, in association with *Pseudomonas* sp. and *Micrococcus* sp., are able to reduce the concentration of DT to 0.01 mg/l and below. However, all other strains studied did not show significant destructive activity in relation to DT.

It is known that simple unbranched n-alkanes predominate in DT (Heath et al., 1993). Although the metabolism of n-alkanes (C6-C12) is possible, many of them act as solvents, destroying cells through partial solubilization of membrane phospholipids, thus exerting a toxic effect on many microorganisms (Cunningham and Philp, 2000). It is possible that the bacteria studied also turned out to be susceptible to the toxic effect of low molecular weight alkanes of diesel fuel.

The results of a quantitative analysis of mineral oil degradation showed that in the samples with *B. firmus* OBR 3.1 and *B. circulans* NSh, a decrease in hydrocarbons was observed, by 39% and 35%, respectively.

These results coincided with the high destructive activity of these cultures in relation to oil. Cultivation of other microbial strains did not lead to significant degradation of mineral oil. The low destructive potential of degrading bacteria in relation to the studied mineral oil (Luxe standard 15W-40) can be associated with the multicomponent composition of this oil product, which, in addition to high-molecular hydrocarbons, includes various additives, such as phenolates, dithiophosphates and alkyl dithiophosphates of various metals (Karaulov and Khudoliy, 2000).

**CONCLUSIONS**

Thus, the experiment showed that all bacteria isolated from BS are good at assimilating oil as the only source of carbon and energy, which may be related to the sources of their release (oil wells) and, as a result, their long-term adaptation to this substrate as a source of nutrition. The destructive activity of the isolated bacteria in relation to diesel fuel and mineral oil was significantly lower, which is consistent with the literature data, because it is

Conclusions deleted are from other authors.

Only your results should be concluded.
known that despite the lower content of aromatic hydrocarbons in their composition, these petroleum products have limited bioavailability for degrading microorganisms (Zanaroli et al., 2010).

The fact that not all cultures growing on a solid medium with mineral oil and diesel fuel showed noticeable activity in suspension culture may also be due to the well-known effect when cells attached to surfaces are more active than free cells (Ananko et al., 2005). It was shown that the cells of a number of strains are unable to assimilate toxic substrates when grown on diffuse agar, but utilize the substrates in agar granules (Fedorov et al., 2000). Such strains are of interest for use in immobilized form.

On the basis of the results of the conducted experiment, the following bacteria can be identified as promising decomposers of petroleum hydrocarbons: B. circulans NSh degrades oil, diesel fuel and mineral oil in 10 days of cultivation in a liquid medium – by 38, 25 and 35%, respectively; B. firmus OBR 3.1 – oil and mineral oil – by 38 and 39%; Halomonas sp. OBR 1 – oil – by 40%; B. circulans OBR 3.3 – DT – by 35.5%.

The fact that not all bacteria isolated from BS showed a high destructive activity with respect to oil products can be explained by the following. It is known that in water-flooded oil fields, where the only source of carbon is oil, along with oil-oxidizing microorganisms ("producers"), many chemoheterotrophic aerobic and anaerobic satellite microorganisms (dissipotrophs) can be distinguished that are not able to utilize oil components, but grow at the expense of products its degradation. Perhaps, among the bacteria isolated, there were just such satellite microorganisms. The authors (Milekhina et al., 1998) noted that the hydrocarbon-oxidizing activity was characteristic only of the gram-positive bacteria isolated from formation fluids and oil from the Bondyuzhskoye and Romashkinskoye oil fields of Tatarstan. On the basis of their results, the researchers suggested that Gram-negative bacteria have a different functional role in oil-bearing ecosystems.

Thus, as a result of the studies carried out, it was shown that BSs are a source of specialized microorganisms capable of decomposing various xenobiotics. At the same time, HOM dominate in microbial communities of BS. A wide substrate spectrum for petroleum hydrocarbons found in isolated bacteria indicates the prospects for using such microorganisms in technologies for microbial utilization of drilling waste.

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