SOME CORROSIVE BACTERIA ISOLATED FROM THE TECHNOGENIC SOIL ECOSYSTEM IN CHERNIHIV CITY (UKRAINE)

Nataliia Tkachuk1*, Liubov Zelena2

1 Department of Biology, T.H. Shevchenko National University “Chernihiv Colehium”, Hetman Polubotok Str. 53, 14013 Chernihiv, Ukraine; e-mail: nataliia.smykun@gmail.com
2 Department of Physiology of Industrial Microorganisms, Danylo Zabolotny Institute of Microbiology and Virology of the National Academy of Sciences of Ukraine, Acad. Zabolotny Str. 154, 03143 Kyiv, Ukraine; e-mail: zelenalyubov@gmail.com

* corresponding author

Abstract:
The soil microbiome is exposed to technogenic influence during the operation of metal structures. There are quantitative and qualitative changes in the microbiota of the technogenic ecosystem. During the study of the technogenic soil ecosystem (ferrosphere), samples of which were taken in the field (Chernihiv, Ukraine: 51°29’58”N, 31°16’09”E), the presence of corrosively active microbial cenosis was established: sulfate-reducing, denitrifying, iron-reducing (using acetate as the only electron donor, and Fe (III) as the only electron acceptor) and ammonifying bacteria. The predominant representatives of corrosively active groups of bacteria were isolated. They were identified as Bacillus simplex, Streptomyces gardneri, Streptomyces canus (ammonifying bacteria), Fictibacillus sp. (ammonifying bacteria with iron-reducing ability), Anaerotignum (Clostridium) propionicum (organic acid-producing bacteria), Desulfovibrio oryzae (sulfate-reducing bacteria) based on some microbiological, physiological and biochemical, genetic features. Strains of heterotrophic and hemolitotrophic bacteria (individual representatives and their associations) isolated from the technogenic ecosystem can be used in both industrial and technological spheres. The interaction of isolated bacteria in the process of microbial induced corrosion is a prospect for further research.

Key words: 16S rRNA gene, microbial induced corrosion, phenotypic characteristics.

Manuscript received 25 November 2020, accepted 17 May 2021

INTRODUCTION

Microbial induced corrosion (MIC) is the result of the activity of microorganisms in the form of a biofilm on a metal or another corroding surface. The development of corrosive groups occurs in the technogenic ecosystem – the ferrosphere, which is the area of soil that comes into contact with metal surfaces (Antonovskaya et al., 1986; Costerton et al., 1995; James et al., 1995; Lewandowski, 2000; Pilyashenko-Novokhatny, 2000; Andreyuk et al., 2002, 2005; Purish and Asaulenko, 2007; Purish et al., 2009). The taxonomic composition is represented by bacteria of the genera Pseudomonas, Arthrobacter, Bacillus, Desulfovibrio, Streptomyces, Thioacillus (Ait-Langoma zino et al., 1991). It has been experimentally proved that changes occur in the microbial group during the formation of a steel biofilm on the surface (Purish and Asaulenko, 2007; Purish et al., 2009). In the first stages of biofilm formation, the dominant position is occupied by heterotrophic denitrifying (DNB) and ammonifying bacteria (AMB), which contribute to the creation of anaerobic conditions on the steel surface. Next sulfate-reducing (SRB) and iron-reducing bacteria (IRB) develop in the biofilm (Purish and Asaulenko, 2007; Purish et al., 2009). The dominance of the previous physiological group creates optimal conditions for the functioning of the next. It is shown that in the case of metal corrosion, gram-negative bacteria are 80% more dominant than gram-positive among the isolated heterotro phic bacteria. The overall distribution was Pseudomonas sp. (20%), Bacillus sp. (10%), Gallionella sp. (10%) and Vibrio sp. (10%) (Nuñez, 2007).

It has become possible recently to define a wider range of microbial variety in natural environments using molecular-genetic approaches (Stahl et al., 1984; Giovannoni et al., 1990; Amann et al., 2002). Today, the number of reports of molecular genetic research in areas with corrosion is growing. In particular, corrosion-active microbial groups of nuclear waste storage (Horn et al., 2003; Engel et al.,
2019), sewage systems (Vincke et al., 2001; Okabe et al., 2007; Satoh et al., 2009), copper pipes (Pavissich et al., 2010), pipelines, seal rings, steel plates, stainless steel coupons in seawater (Lopez et al., 2006; Bermont-Bouis et al., 2007; Li et al., 2017; Capão et al., 2020; Salgar-Chaparro et al., 2020), gas pipelines (Zhu et al., 2003; Jan-Roblero et al., 2004), oil pipelines (Magot et al., 1997; Neria-Gonzalez et al., 2006; Su et al., 2019), galvanized steel pipes (Bolton et al., 2010) have been investigated. A higher complexity of microbial groups than observed with bacterial culture approaches was found out. However, no common microbial groups or general trends were observed in these studies, as the areas of corrosion studied were diverse and complex.

The aim of this study was to isolate the predominant representatives of ammonifying, iron-reducing, organic acid-producing and sulfate-reducing bacteria from the technogenic soil ecosystem (ferrosphere), and to identify them using microbiological and molecular-genetic methods.

MATERIAL AND METHODS

Sampling

A soil selection of the technogenic ecosystem (ferrosphere) was carried out in the field in November from the surface of a metal structure (fence support, 15 years immersed in sod-podzolic soil) from generally accepted sampling depth 0.7 m in studies of the sulfidogenic microbial community (Abdulina et al., 2011). Sampling site is located in Chernihiv (Ukraine), 51°29’58”N 31°16’09”E (Fig. 1). The sample was collected using a sterile glass flask and was immediately transported to the laboratory of T.H. Shevchenko National University “Chernihiv Colehium” and stored at 4°C in the refrigerator. Further definition of the determination of number of bacteria of corrosive dangerous groups, the isolation of bacteria and the studies of cultural, morphological, physiological and biochemical properties of isolates were carried out in T.H. Shevchenko National University “Chernihiv Colehium”.

Isolation of the strains of bacteria and identification

The isolation of bacteria was carried out from dilution 10^{-1}–10^{-6} (depending on the number of microorganisms of the corresponding corrosive group) on MPA (ammonifying bacteria – ChNPU F1, ChNPU F3, NUChC F2 isolates), FWA-Fe (III) citrate medium (iron-reducing bacteria – ChNPU ZVB1 isolate), Postgate’s “B” medium (organic acid-producing bacteria (NUChC Sat1 isolate) and sulfate-reducing bacteria (NUChC SRB1, NUChC SRB2 isolates). The purity of the culture was verified by microscopy and cultivation on various nutrient media (potato agar, oat agar, peptone-yeast agar, Chapek’s medium, starch-ammonia agar). An optical microscope Delta Optical Genetic Pro with increase ×400 and ×1000 and electron microscope BS-540 (Tesla, Czechoslovakia) with magnification x16000 were used to study the morphology of bacteria. Determination of the size of bacterial cells was carried out using the Aigo Scopelmage 9.0 computer program.

The studies of cultural, morphological, physiological and biochemical properties of isolated bacteria were carried out by the conventional methods (Methods of general bacteriology, 1984). Methods of molecular genetic study of bacteria (sequence of 16S rRNA gene fragments and phylogenetic analysis) were also used by the procedures which were described by Tkachuk et al. (2017) and were carried out in Danylo Zabolotny Institute of Microbiology and Virology of the National Academy of Sciences of Ukraine. Briefly, to perform 16S rRNA gene sequencing genomic DNA was isolated from pure bacterial cultures and amplification with 27F and 1492r primers (Lane, 1991) was carried out. PCR-products were purified and sequenced in both directions using ABI 310 (Applied Biosystems). Obtained sequences were compared with those deposited in GenBank database with BLASTn and the highest revealed percent of similarity between sequences (>98.7%) was taken to

![Fig. 1. Location of sampling site.](image-url)
identify species. In case of multiple similarity sequences with >98.7% were searched and analyzed for variable and parsimony-informative sites with MEGA6 (Tamura et al., 2003) to specify identification.

Statistical analysis of experimental data

Methods of mathematical statistics were used in processing the data on the number of microorganisms (Plohinskij, 1970). The number of microorganisms on liquid media was determined using McCready tables and on solid media – taking into account the mean square deviation.

RESULTS

Microbiological analysis of the soil from ferrosphere allowed us to determine a number of microorganisms of corrosion-active groups. It has been found out that the number of sulfate-reducing bacteria in the investigated soil is quite high – 6.6×10^4 cells/1 g of abs. dry soil. The number of denitrifying bacteria in the soil of the ferrosphere appeared to be similar with SRB (6.6×10^4 cells/1 g of abs. dry soil). However, the number of iron-reducing bacteria in the studied soil was insignificant including 10^3 cells/1 g of abs. dry soil. The number of heterotrophic ammonifying bacteria in the soil of the ferrosphere is 1.19±0.03×10^6 of colony-forming units/1 g of abs. dry soil.

Thus, the number of microorganisms of corrosive dangerous groups in the studied technogenic soil indicates presence of corrosively active microbial cenosis.

Next, we isolated the predominant representatives of corrosive dangerous groups of bacteria from the soil technogenic ecosystem. The results of the study of cultural-morphological and physiological-biochemical properties of the isolated bacteria are presented (Tables 1–7).

Microbiological characteristics according to Bergey’s Manual of Systematic Bacteriology (Bergey’s Manual of Systematic Bacteriology, 2005, 2009, 2012) prove that the isolated bacteria can belong to the Clostridiaceae family (NUChC Sat1 isolate), the Streptomycetes family (NUChC SRB1, NUChC SRB2 isolates), the Clostridiales family (NUChC F2 isolate), the Anaerobacteriales family (NUChC Sat1 isolate), the Deferribacteraceae family (NUChC F2 isolate), the Desulfovibrionaceae family (NUChC SRB1, NUChC SRB2 isolates).

To define species of isolates 16S rRNA, the gene sequencing was performed. As a result nucleotide sequences of gene fragments were obtained and compared to those deposited in GenBank database (Table 8). Generally, the length of 16S rRNA gene fragment varied from 483 bp to 1368 bp and was informative for genus or species, in most cases, identification. The phylogenetic analysis including detailed nucleotide composition analysis and construction of phylogenetic dendrogram was carried out where the defining of species was controversially.

Based on microbiological, physiological and biochemical, genetic features, the investigated bacteria are identified as Bacillus simplex, Streptomyces gardneri, Streptomyces canus, Fictibacillus sp., Anaerotignum (Clostridium) propionicum and Desulfovibrio oryzae (Table 8).

DISCUSSION

Bacteria of the Bacillus genus deserve attention as one of the representative microorganisms for MIC research (Wang et al., 2020). There are no reports of involvement of B. simplex and Fictibacillus sp. in microbial damage to materials. However, according to the analyzed publications, microbial damage to metals depends on the studied species and strain of bacteria of the Bacillus genus (Rajasekar and Ting, 2010; Bano and Qazi, 2011; Bragadeeswaran et al., 2011; Du et al., 2014). The metabolites of Bacillus sp. contain cyclic peptides, antibiotics, polyglutamic acid, polyaspartic acid and poly polysaccharide, and can form biofilms with spores. Research showed that extracellular polymeric substances produced by Bacillus was the main component of the biofilm formed on the metal surface and provided the biologically active compound which affected the corrosion process. More and more attention has been paid to the corrosion effect of Bacillus on metals, but the mechanism is still unclear, and there is still much controversy (Wang et al., 2020).

Streptomycetes of isolated species (S. gardneri, S. canus) are also not known to participate in the biodamage of materials. However, the ability to produce biologically active substances known to streptomycetes determines the interest in microbial corrosion studies involving these bacteria (Jayaraman et al., 1997; Nnabuk Eddy Okon, 2010; Pacheco da Rosa et al., 2013, 2016; Bleich et al., 2015).

The D. oryzae species is currently poorly studied. In the available literature there is a single brief description of the species only (Salgar-Chaparro and Silva-Plata, 2008). Some D. oryzae strains have been isolated from a community of oil-related water associated with the corrosion process (Salgar-Chaparro and Silva-Plata, 2008; Duque et al., 2011).

Among bacteria, secreting organic acids and exopolymers or slime clostridia deserve attention. They have been isolated from the sites of corrosion (Pope et al., 1984; Beech and Gaylarde, 1999; Zha et al., 2003; Oliveira et al., 2011), sulfidogenic bacterial consortia (Herro and Port, 1993; Agrawal et al., 2010; Monroy et al., 2011; Ilhan-Sungur et al., 2017; Qiu et al., 2017). Herewith, the mechanisms of influence on the process of MIC and the consequences depend on the species of clostridia. In general, the corrosion hazard for metal coatings reveals the products of clostridium metabolism (organic acids, slim, H2, H2S), followed by synergism between Clostridium sp. and SRB, which helps to increase the corrosive aggressiveness of the environment (Beech and Gaylarde, 1999; Zha et al., 2003; AlAbbas et al., 2013; Bala and Chidambaram, 2014). Co-cultivation of D. oryzae with A. propionicum (without electron donors, the presence of sulfates and yeast extract) showed the formation of hydrogen sulfide by sulfate-reducing bacteria, which was not observed in the case of cultivation of monocultures (Tkachuk et al., 2020). In this case, a phenomenon
Table 1. Cultural and morphological properties of ChNPU F1 isolate.

<table>
<thead>
<tr>
<th>Sign</th>
<th>Characteristic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Colonies on the MPA</td>
<td>Colonies are superficial, grayish-white, 1–2 mm in diameter (1st day) and 3–6 mm (2nd days). The shape of the colonies is rounded, the edges are uneven, wavy. The profile is convex, the structure is homogeneous, the surface is smooth, shiny. The texture is soft. Colonies are translucent, opalescent in transmitted light after 18 hours of cultivation. On the 2nd day the colonies are opaque. On the 7th day the colonies become concentric, the profile acquires a conical shape.</td>
</tr>
<tr>
<td>Growth in the MPB</td>
<td>The solution is moderately uniformly turbid, the film is thin, smooth; no smell; no pigmentation; sediment is significant, soft</td>
</tr>
<tr>
<td>Colonies on potato agar</td>
<td>The isolate grows on the 1st day. The shape of the colonies is rounded, grayish-white with opalescent wavy edges. The diameter of the colonies is 2–3 mm on the 1st day and 15–18 mm on the 3rd day. The profile of the colonies is flattened, crater-like on the 3rd day, the structure is homogeneous, the surface is smooth and shiny. The texture is soft.</td>
</tr>
<tr>
<td>Colonies on peptone-yeast agar</td>
<td>The isolate grows on the 1st day. Colonies are with a diameter of 0.1 mm (1st day) and 3–5 mm (3rd day), whitish-gray, round with jagged edges. The structure of the colonies is fine-grained, the profile is flattened, the surface is uneven, folded, matte. The texture is soft.</td>
</tr>
<tr>
<td>Colonies on Chapek’s medium</td>
<td>The isolate grows on the 3rd day. Colonies are small (diameter 0.1 mm), grayish-white, rounded with jagged, wavy edges. The profile is convex, the surface is smooth, shiny. The texture is soft.</td>
</tr>
<tr>
<td>Colonies on starch-ammonia agar</td>
<td>The isolate grows on the 2nd day. Growth is weak. Colonies are of irregular shape, membranous, 4–5 mm in diameter, white-gray, with uneven, wavy edges. The structure of the colonies is fine-grained, the profile is flattened, the consistency is soft.</td>
</tr>
<tr>
<td>The shape and location of cells</td>
<td>Cells in young culture (18 hours) are rod-shaped, with rounded ends, 1–2 cells are located. In the old culture (2 months) the cells are rod-shaped with rounded ends, 1–2 cells are located, in chains of 4 or more cells.</td>
</tr>
<tr>
<td>Cell length</td>
<td>5.80±0.25 μm</td>
</tr>
<tr>
<td>Cell motility</td>
<td>Weak motility</td>
</tr>
<tr>
<td>Endospores, their location</td>
<td>Available, central location</td>
</tr>
<tr>
<td>Gram stain</td>
<td>Gram-negative</td>
</tr>
</tbody>
</table>

Table 2. Cultural and morphological properties of ChNPU F3 isolate.

<table>
<thead>
<tr>
<th>Sign</th>
<th>Characteristic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Colonies on the MPA</td>
<td>Colonies are superficial, round, 0.5–3 mm in diameter, yellowish-gray with smooth edges. The structure is homogeneous, the profile is convex, the surface is smooth, matte. The consistency is oily. On the 14th–30th days of cultivation, colonies up to 10 mm in diameter have a weak white air mycelium.</td>
</tr>
<tr>
<td>Colonies on oat agar</td>
<td>Colonies are gray, rounded, 0.5–1.0 mm in diameter. The edges are fringed, the surface is matte, the profile is convex, the consistency is firm. Air mycelium is not expressed. The reverse side of the colonies without different pigments (yellowish-gray color). Over time, the diameter of the colonies on oat agar increases (2.5–3.0 mm), the shape becomes rounded with a roller, the profile is crater-like, air mycelium is not expressed. On the 10th day of cultivation, the colonies have white air mycelium, weakly expressed.</td>
</tr>
<tr>
<td>Growth in the MPB</td>
<td>The solution is moderately uniformly turbid, the film is thin, smooth; no smell; no pigmentation; sediment is significant, soft</td>
</tr>
<tr>
<td>Colonies on potato agar</td>
<td>The isolate grows on the 3rd day. Colonies are rounded, folded, with wavy edges, yellowish-white, 2–3 mm in diameter. The profile is convex, the structure is fine-grained, the surface is matte, the texture is soft. When the colony is removed by the bacteriological loop, the substrate mycelium remains. Air mycelium is not expressed. Pigments are not formed.</td>
</tr>
<tr>
<td>Colonies on peptone-yeast agar</td>
<td>The isolate grows on the 3rd day. Colonies with a diameter of 2–3 mm are rounded with wavy edges, white. The surface is folded, matte, flat profile, fine-grained structure, soft texture. There is a substrate mycelium. Air mycelium is not expressed. Pigments are not formed.</td>
</tr>
<tr>
<td>Colonies on Chapek’s medium</td>
<td>The isolate grows on the 3rd day. Colonies with a diameter of 1 mm round shape with fringed edges, white. The surface is matte. The profile is flattened, the structure is homogeneous, the consistency is firm. There is a substrate mycelium. Air mycelium is not expressed. Pigments are not formed.</td>
</tr>
<tr>
<td>Colonies on starch-ammonia agar</td>
<td>The isolate grows on the 2nd day. Colonies are with a diameter of 0.1 mm, rounded, grayish-white, convex profile, fringed edges, fine-grained structure, firm consistency. There is a substrate mycelium. Air mycelium is not expressed.</td>
</tr>
<tr>
<td>The shape and location of cells</td>
<td>Form a branched mycelium with spores of round shape, arranged in a chain of 5–10 spores or more</td>
</tr>
<tr>
<td>Endospores, their location</td>
<td>None</td>
</tr>
<tr>
<td>Gram stain</td>
<td>Gram-positive</td>
</tr>
</tbody>
</table>
### Table 3. Cultural and morphological properties of NUChC F2 isolate.

<table>
<thead>
<tr>
<th>Sign</th>
<th>Characteristic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Colonies on the MPA</td>
<td>The isolate grows on the 4th day, the colonies are rounded, 2–4 mm in diameter, gray-brown, the surface is folded, leathery, matte, coarse-grained structure, firm consistency. Releases a dark brown pigment. Air mycelium is not formed.</td>
</tr>
<tr>
<td>Colonies on oat agar</td>
<td>Colonies are superficial, gray-white, 2–4 mm in diameter. The shape is rounded with a roller, the profile is crater-shaped, the surface is folded, the consistency is fibrous. On the 14th–20th day, a well-defined white air mycelium develops. On the underside of the colony is brown. The isolate forms a brown pigment, which over time (1–2 months) becomes reddish.</td>
</tr>
<tr>
<td>The shape and location of cells</td>
<td>Bacteria form a branched mycelium with spores arranged in a chain of 10–50 spores or more. Spore chains are spiral, but there are some spore chains straight and loop-like.</td>
</tr>
<tr>
<td>Endospores, their location</td>
<td>None</td>
</tr>
<tr>
<td>Gram stain</td>
<td>Gram-positive</td>
</tr>
</tbody>
</table>

### Table 4. Cultural and morphological properties of ChNPU ZVB1 isolate.

<table>
<thead>
<tr>
<th>Sign</th>
<th>Characteristic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Colonies on FWA-Fe (III) citrate medium</td>
<td>Colonies grow on the 7th day under both anaerobic and aerobic cultivation conditions. Colonies are superficial, amber in color, 4 mm in diameter, irregularly shaped, flattened, wavy edges, matte surface, coarse-grained structure, soft texture.</td>
</tr>
<tr>
<td>Colonies on the MPA</td>
<td>Colonies of the isolate grow at 29°C on the 1st day under aerobic conditions. Colonies are superficial, round, yellowish-gray, 1–2 mm in diameter, convex profile, wavy edges, shiny surface, smooth, homogeneous structure, soft texture.</td>
</tr>
<tr>
<td>Growth in the MPB</td>
<td>The solution is moderately uniformly turbid; the film is thin, well defined, smooth; sediment is significant, loose; there is a specific smell; no pigmentation.</td>
</tr>
<tr>
<td>Colonies on potato agar</td>
<td>Colonies grow on the 3rd day, rounded, jagged edges, hairy. The profile is flattened. The surface is shiny. The structure is homogeneous. The texture is soft.</td>
</tr>
<tr>
<td>Colonies on peptone-yeast agar</td>
<td>Colonies grow on the 1st day, small, 2–3 mm in diameter, yellowish-gray color. The shape is rounded. The edges are uneven. The surface is smooth, shiny. The profile is flattened with the acting center. The structure is fine-grained. The texture is soft.</td>
</tr>
<tr>
<td>Colonies on Chapek’s medium</td>
<td>It does not grow</td>
</tr>
<tr>
<td>Colonies on starch-ammonia agar</td>
<td>Weak, membranous growth</td>
</tr>
<tr>
<td>The shape and location of cells</td>
<td>Slightly curved sticks with rounded ends, combine into long curved threads surrounded by a cover, some cells are stretched.</td>
</tr>
<tr>
<td>Cell length</td>
<td>4.355±0.204 μm</td>
</tr>
<tr>
<td>Cell motility</td>
<td>Motility</td>
</tr>
<tr>
<td>Endospores, their location</td>
<td>Available, central location</td>
</tr>
<tr>
<td>Gram stain</td>
<td>Gram-positive</td>
</tr>
</tbody>
</table>

### Table 5. Cultural and morphological properties of NUChC Sat1 isolate.

<table>
<thead>
<tr>
<th>Sign</th>
<th>Characteristic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Colonies on the Postgate’s “B” medium (anaerobic conditions)</td>
<td>Colonies are deep, grayish-beige, round, soft, 2–3 mm in size</td>
</tr>
<tr>
<td>Shape and location of cells</td>
<td>Stick-shaped, slightly curved, with rounded ends, single</td>
</tr>
<tr>
<td>Cell length</td>
<td>4.671±0.387 μm</td>
</tr>
<tr>
<td>Cell motility</td>
<td>Motility</td>
</tr>
<tr>
<td>Endospores, their location</td>
<td>Spores are sub-terminal and inflate cells</td>
</tr>
<tr>
<td>Gram stain</td>
<td>Gram-positive</td>
</tr>
</tbody>
</table>

### Table 6. Cultural and morphological properties of NUChC SRB1 and NUChC SRB2 isolates.

<table>
<thead>
<tr>
<th>Sign</th>
<th>Characteristic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Colonies on the Postgate’s “B” medium (anaerobic conditions)</td>
<td>Clear black SRB colonies with a diameter of 1 mm (NUChC SRB1) and 2 mm (NUChC SRB2)</td>
</tr>
<tr>
<td>The shape and location of cells</td>
<td>Vibrios with rounded ends, respectively, monotrichs</td>
</tr>
<tr>
<td>Cell length</td>
<td>4.6±0.3 μm (NUChC SRB1), 1.8±0.1 μm (NUChC SRB2)</td>
</tr>
<tr>
<td>Cell motility</td>
<td>Motile</td>
</tr>
<tr>
<td>Endospores, their location</td>
<td>No sign</td>
</tr>
<tr>
<td>Gram stain</td>
<td>Gram-negative</td>
</tr>
</tbody>
</table>
Table 7. Some physiological and biochemical properties of isolated bacteria.

<table>
<thead>
<tr>
<th>Isolate</th>
<th>Physiological and biochemical properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>ChNPU F1</td>
<td>Catalase-positive; oxidase-negative; aerobic; citrate utilize; do not form indole; formic acid fermentation is not carried out; casein is not hydrolyzed; do not form hydrogen sulfide and cellulases; ammonia is formed</td>
</tr>
<tr>
<td>ChNPU F3</td>
<td>Catalase-positive; oxidase-negative; aerobic; do not utilize citrate; utilize urea; do not form indole; formic acid fermentation is not carried out; levan is not synthesized; starch, fats, casein, gelatin are hydrolyzed; do not form hydrogen sulfide and cellulases; ammonia is formed</td>
</tr>
<tr>
<td>NUChC F2</td>
<td>Catalase-positive, oxidase-negative; aerobic, do not utilize glucose, but grow due to the amino acids peptone, releasing ammonia; hydrolyze starch and casein; do not hydrolyze lipids and urea; citrate is utilized; do not break down aromatic amino acids; do not form hydrogen sulfide; ammonia is produced; levan saccharase is absent; formic acid fermentation is not carried out, Fe (III) is reduced</td>
</tr>
<tr>
<td>ChNPU ZVB1</td>
<td>Catalase and oxidase-positive, facultative anaerobes, glucose is utilized both under aerobic conditions (with the formation of acid but not gas) and under anaerobic conditions (with the formation of acid but not gas); citrate urea is not utilized; do not form indole; formic acid fermentation is not carried out; levan is synthesized; fats and gelatin are hydrolyzed; starch, casein are not hydrolyzed; have the phospholipase; do not grow on manniot salt agar, which may be due to sensitivity to high concentrations of NaCl; form ammonia; do not form hydrogen sulfide and cellulase</td>
</tr>
<tr>
<td>NUChC Sat1</td>
<td>Catalase- and oxidase-negative; the isolate assimilates lactate, but does not assimilate glycerol, dulcite, ethanol, mannitol, sorbitol, arabinose, lactose, ramnose, raffinose, starch, sucrose as a source of Carbon.</td>
</tr>
<tr>
<td>NUChC SRB1,NUChC SRB2</td>
<td>Catalase-negative, oxidase-positive; able to use lactate and formate as electron donors. Acetate, fumarate, propionate, malate, fructose and glucose are not used as electron donors; NUChC SRB1 isolate uses sulfate and thiosulfate as electron acceptors, but does not use sulfate, fumarate and nitrate. NUChC SRB2 isolate uses sulfate, thiosulfate and fumarate as electron acceptors, but does not use sulfate and nitrate.</td>
</tr>
</tbody>
</table>

Table 8. Ecological and trophic group, the number of the registration of nucleotide sequences of isolated and identified bacteria.

<table>
<thead>
<tr>
<th>Species or genus</th>
<th>Ecological and trophic group</th>
<th>Strain</th>
<th>Number of registration in the GenBank</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacillus simplex</td>
<td>ammonifying bacteria</td>
<td>ChNPU F1</td>
<td>KX349220</td>
</tr>
<tr>
<td>Streptomyces gardneri</td>
<td>ammonifying bacteria</td>
<td>ChNPU F3</td>
<td>KX349221</td>
</tr>
<tr>
<td>Streptomyces canus</td>
<td>ammonifying bacteria</td>
<td>NUChC F2</td>
<td>MG924748 and MG924855</td>
</tr>
<tr>
<td>Fictibacillus sp.</td>
<td>ammonifying and iron-reducing bacteria</td>
<td>ChNPU ZVB1</td>
<td>KX349222</td>
</tr>
<tr>
<td>Anaerotignum (Clostridium) propionicum</td>
<td>organic acid-producing bacteria</td>
<td>NUChC Sat1</td>
<td>MG924854.1</td>
</tr>
<tr>
<td>Desulfovibrio oryzae</td>
<td>sulfate-reducing bacteria</td>
<td>NUChC SRB1</td>
<td>MT102713.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NUChC SRB2</td>
<td>MT102714.1</td>
</tr>
</tbody>
</table>

of syntrophy seems to take place and the electron donor appears due to the use of the yeast extract compounds by the NUChC Sat1 strain.

CONCLUSIONS

Thus, the soil microbiome is exposed to technogenic influence during the operation of metal structures in it, there are quantitative and qualitative changes in the microflora. At the same time, strains of heterotrophic and heliotrophic bacteria (individual representatives and their associations) isolated from the technogenic ecosystem can be used in both industrial and technological spheres. The interaction of isolated bacteria in the process of microbial induced corrosion is a prospect for further research.

Acknowledgements

We are sincerely grateful to the leading engineer Volodymyr Strekalov (Institute of Agricultural Mnmiobiology and Agroindustrial Production of the National Academy of Agricultural Sciences of Ukraine) for carrying out the electronic microscopy.

REFERENCES


CORROSIVE BACTERIA FROM THE TECHNOGENIC SOIL ECOSYSTEM


