Impact of Algicidal *Bacillus mycoides* on Diatom *Ulnaria acus* from Lake Baikal

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Abstract: Algae–bacteria interactions play an important role in water ecosystems. In this work, the BS2-15 algicidal strain was isolated from the bottom sediments of Lake Baikal and identified as *Bacillus mycoides* on the basis of 16S rDNA sequencing, its described ultrastructure, and biochemical properties. *B. mycoides* BS2-15 was demonstrated to have a strong algicidal effect against a freshwater diatom culture of *Ulnaria acus*, inhibiting its growth and increasing frustules fragility. By analyzing the impact of bacterial filtrate onto the cells of *U. acus*, we demonstrated that perhaps an algicidal compound is produced by bacteria independently in the presence of diatoms in a medium. Using methods of TUNEL and confocal microscopy, we revealed that the bacterial algicidal effect on the diatom cells results in DNA fragmentation, nucleus destruction, and neutral lipid accumulation. This phenomenon highlights the complexity of algae–bacteria interactions and their potential role in regulating water ecosystem microbial populations.

Keywords: diatoms; algicidal bacteria; *Bacillus mycoides*; TUNEL

1. Introduction

Phytoplankton is essential for global carbon cycles [1], and diatom algae in particular have an organic carbon output comparable to that of land plants [2,3]. Primary producers in water ecosystems have complex relationships with bacteria [4]. Along with the search for algicidal bacteria useful in preventing toxic algal blooms [5], the mechanisms of algae–bacteria interactions have fundamental importance [6]; many bacterial strains have an algicidal effect on diatoms by releasing enzymes into the environment [7–9].

*Bacillus* spp. are Gram-positive bacteria widely distributed in the environment; this is likely because they form endospores resistant to different factors, such as drying out, UV radiation, and lack of nutrients [10,11]. *Bacillus mycoides* is characterized by its rhizoidal growth in agar medium [12]. In addition, *B. mycoides*, like many representatives of the genus *Bacillus*, are saprophytic organisms [13,14].

The nature of algicidal compounds produced by bacteria into the medium can be very diverse. There are many known algicidal secondary metabolites, which can be produced by bacteria of the genus *Bacillus* into the environment [15–18]. Moreover, it was demonstrated that *B. mycoides* negatively affect pathogenic fungi, as it emits toxic volatile compounds ammonia and dimethyl sulfide [19]. However, cytological manifestations of the algicidal effect have not been described. In this work, we studied the algicidal effect of *B. mycoides* BS2-15, a strain isolated from sediments of Lake Baikal.

The diatom *Ulnaria acus* (=*Synedra acus*) is a member of the dominant assemblage of recent Baikal phytoplankton, but its remains in the upper layer of bottom sediments and cannot be found in all regions of the lake because it is subject to considerable degradation,
both in the water column and in the surface sediment layer [20]. The axenic monoclonal culture *U. acus* BK280 was applied as a model for multiple experimental studies [20–26].

Sediments of Lake Baikal were earlier found to contain bacteria involved in the destruction of diatom cells [9]. The bacteria *B. mycoides* BS2-15 and diatom *U. acus* were isolated from Lake Baikal. The aim of the study was to determine the effect of the algicidal impact of *B. mycoides* BS2-15 on *U. acus* BK280 and to reveal cytological manifestations of the diatom cells.

2. Materials and Methods

2.1. Diatom Culture

A culture of diatom microalgae *U. acus* was grown in March 2013 from a subglacial sample taken at Lake Baikal near Bolshye Koty settlement. Monoclonal diatom cultures were grown in Diatom medium (DM) [27] at 10 °C with illumination of 16 µmol/m²/s under a 12:12 day/light cycle. An axenic culture of *U. acus* BK280 was produced according to a published protocol [28]. The procedure involves filtration through a polycarbonate membrane with a pore size of 5 µm (to remove free bacteria), detergent treatment (20 µg/mL Triton X-100), cell treatment with the selected antibiotic (5 µg/mL ciprofloxacin, 18 h incubation), repeated filtration, and monoclonal culturing of diatom cells. Axenity of the culture was verified microscopically (after DAPI staining) and by molecular biological methods. The diatom cells were incubated and transferred to fresh DM media once every four weeks.

2.2. Isolation of Algicidal Bacteria

Sediment samples were taken using a benthic gravity corer aboard the research vessel ‘Vereschagin’. Sampling took place in southern Baikal (51°47′44″ N, 104°54′26″ E) in August 2010. The upper 10 cm of the sediment core was immediately sliced into 1 cm layers. The sediment from each layer (0.5 g) was mortared, placed in 100 mL of sterile distilled water and shaken for two hours. After settling for ten minutes, 1 mL of supernatant was spread on Petri dishes with tenfold diluted fish peptone agar (FPA/10) and cultivated under aerobic conditions at 25 °C for seven days. Pure bacterial cultures were isolated on the same medium using the streak inoculation technique. The morphological properties of the strains were examined using optical microscopy (Axiovert 200, Zeiss, Germany), and the physiological and biochemical properties and taxonomic identity were investigated as described previously [21].

2.3. Identification of Selected Bacterium

The identification of strain BS2-15 was based on its 16S ribosomal RNA sequence. A phenol-chloroform extraction protocol was applied for DNA purification, using lysozyme (50 mg/mL) and 10% SDS [20,29]. DNA was amplified using a PCR kit (Amplisens, Moscow, Russia) and universal eubacterial 16S rRNA primers: 27L (5′-AGAGT TTGATC ATGCTCAG-3′) and 1350R (5′-GACGGGCGGTGTGTA CAG-3′) in a BIS M-111 automatic thermocycler (BIS-N, Koltsovo, Russia) under the following conditions: 94 °C for 2 min; 94 °C for 30 s, 55 °C for 40 s, 72 °C for 50 s, 25 cycles; 72 °C for 2 min. The PCR products were analyzed in 1% agarose gel and purified with a CleanupStandard kit (Eurogen, Moscow, Russia). Sequencing was performed with a BigDye V3.1 Terminator Cycles Sequencing Kit and AmpliTaq DNA polymerase FS (Applied Biosystems, Waltham, USA) on an ABI 3130XL Genetic Analyser sequencer at the SB RAS Genomics Core Facility (Novosibirsk, Russia). Analysis of nucleotide sequences of the 16S ribosomal RNA gene was performed by searching reference databases (Genbank) for similar sequences using BLASTN [30]. MEGA 7.0 (Nearest-Neighbor-Interchange (NNI)) was used for phylogenetic analysis [31]. The 16S rDNA sequence of strain BS2-15 was deposited in the GenBank database under accession number MH638320.1.
2.4. Determination of Algicidal Activity

For all the experiments, the cells of BS2-15 from the medium FPA/10 were subcultured into 1% peptone solution and cultured at 25 °C for two days and nights until they reached an optical density of 0.9–1.0 at A600 nm. To determine the dose-dependency of the algicidal effect of bacteria, the cells of *B. mycoides* were added at dilutions 2, 10, 40, 60, 100, 200, 300 µL per mL of medium to the culture of *U. acus* at the exponential phase (for 7 days of culturing, 10⁵ cells/mL) and were co-cultivated for 17 days. To further investigate the algicidal effect, the bacterial inoculum was added to the cells at a diluted 100 µL/mL. To determine if an algicidal agent is produced in the environment regardless of the presence of diatoms near bacteria, cell-free bacterial filtrate (100 µL/mL) ion obtained by filtration of bacterial culture through a sterile 0.20 µm filter (Whatman, Corning, USA), was added to the cells of *U. acus*. To determine whether direct interaction between bacteria and diatoms is necessary for an algicidal effect, bacterial cells washed from the medium were added to the cells at a dilution of 100 µL/mL; they were washed twice with DM medium with precipitation by centrifuge in a sterile test tube at 5000 rpm for 10 min. To exclude the suppressing effect of peptone on the growth of diatom algae, 1% sterile peptone solution was added to diatom culture diluted 100 µL/mL. The cultivation lasted nine days.

For all experiments, we used three biological and three technical replicates. Cells were counted every two days at the same time. An optical and epifluorescent microscope (Axiovert 200 Zeiss, Oberkochen, Germany) was used to enumerate diatom cells and identify bacteria, as previously described [20].

2.5. TUNNEL Assay

Fragmentation of genomic DNA was detected using dUTP end-labelling in situ (TUNEL). Axenic culture *U. acus* cultured with bacteria (100 µL/mL) for 24 h were used for this analysis. Cells were fixed with 3.7% formaldehyde solution (Sigma-Aldrich, St. Louis, MO, USA) with 0.1 M phosphate buffer (pH 7.4) for 15 min. After washing with the phosphate buffer, cells were permeabilized with 0.25% Triton™ X-100 (Sigma-Aldrich, St. Louis, USA). Detection of fragmented DNA was performed with a Click-iT™ TUNEL Alexa Fluor™ 488 Imaging Assay (Thermo Fischer Scientific, Waltham, MA, USA) according to the manufacturer’s protocol. The positive control sample was treated with DNase I (Thermo Fischer Scientific, USA) at room temperature for 30 min before the labeling reaction. Cell nuclei were stained with Hoechst 33,342 (Component F, Click-iT™ TUNEL Alexa Fluor™ 488 Imaging Assay). Prepared cells were placed in a ProLong® Gold Antifade Mountant (Thermo Fisher Scientific, USA). The samples were examined using a Zeiss LSM 710 confocal microscope (Zeiss, Oberkochen, Germany) with a Plan-Apochromat 63×/1.40 Oil DIC M27 objective. Hoechst 33,342 dye was excited at a wavelength of 405 nm and its emission registered between 410–483 nm. Click-iT™ TUNEL Alexa Fluor™ 488 dye was excited at 488 nm and was registered between 493–628 nm.

2.6. Lipid Droplets Analysis Using BODIPY

To determine lipid accumulation used diatom cells, on the seventh day of cultivation axenic culture and with bacterial culture, bacterial filtrate and peptone (all of the above was added to the culture of diatoms at a diluted 100 µL/mL). Diatoms cells were stained with BODIPY 493/503 (Thermo Fischer Scientific, Waltham, USA) by adding the dye to 500 µL of the cells suspension at a concentration of 2 µM and incubating it for 15 min. Then the cells were washed with 0.1 M phosphate buffer (pH 7.4) and fixed with 4% formaldehyde solution; subsequently, they were washed twice with 0.1 M phosphate buffer. Prepared cells were placed in a ProLong® Gold Antifade Mountant (Sigma-Aldrich, St. Louis, USA).

2.7. Fluorescent Microscopy

The cells of axenic culture, when cultured with bacteria and with peptone (both of which are added at a dilution of 100 µL/mL) were stained with the fluorescent dye Live Cell Labeling Kit (Abcam, Cambridge, UK). First 200 µL of the culture aliquot was added
to a 96-well plate. The dye solution was added at a dilution of 0.5 µL/mL, and incubation was performed for 30 min at 10 °C with an illumination of 16 µmol/m²/s. To determine the proportion of living cells in the culture, the stained cells were counted among 100 randomly selected cells using epifluorescence microscope Axiovert 200 (Zeiss, Oberkochen, Germany) with a dark-blue light filter. Staining and counting were performed every two days for 9 days, all counts were performed in triplicate.

2.8. Transmission Electron Microscopy

Diatom cells were fixed with 2.5% glutaraldehyde (Sigma-Aldrich, St. Louis, USA) in 0.1 M phosphate buffer, pH 7.4, for 2 h and postfixed with 1% OsO₄ (Sigma-Aldrich, St. Louis, USA) in the same buffer for 2 h at room temperature. Thereafter, cells were dehydrated in an ascending ethanol series followed by 96% ethanol and acetone dehydrated with copper sulfate for 5 min each (all reagents were obtained from Reakhimkomplekt, Russia). Dehydrated samples were embedded with mixtures of Araldite 502 epoxy resin (Sigma-Aldrich, St. Louis, USA) and acetone and with pure Araldite, transferred to a new portion of Araldite supplemented with DMP-30 accelerator (Sigma-Aldrich, St. Louis, USA), and polymerized at 60 °C for three days. Ultrathin sections were prepared using an Ultracut R ultratome (Leica, Wetzlar, Germany) with an ULTRA 35 diamond knife (Diatom, Nidau, Switzerland), placed onto copper grids, and contrasted with uranyl acetate and lead citrate. TEM analysis was performed using a Leo 906 E microscope (Zeiss, Oberkochen, Germany) at an acceleration voltage of 80 kV. Microscopic images were taken with a MegaView II camera (Zeiss, Oberkochen, Germany) and processed using the MegaVision program.

2.9. Scanning Electron Microscopy

The cell suspension was dehydrated in an ascending ethanol series followed by 96% ethanol and then placed on SEM stubs. These plates were dried at room temperature and coated with gold. Microscopy was performed using a Quanta 200 scanning electron microscope (FEI Company, Hillsboro, OR, USA).

2.10. Statistical Analysis

Statistical assessment of the significance of the differences between retrievals was performed using the program Past 4 with a nonparametric Mann–Whitney U-test, taking into account the retrieval size and normality determined by the Shapiro-Wilk criterion. The differences were considered statistically significant at \( p < 0.05 \).

3. Results and Discussion

3.1. Isolation and Identification of Algicidal Bacterium

We isolated 40 bacterial strains from the bottom sediments of Lake Baikal and assessed for their algicidal effect on the diatom algae U. acus BK280. Eight strains depressed the growth of a monoclonal axenic culture of this alga. Strain BS2-15 (isolated from the upper layer of bottom sediments, 2 cm above surface sediments) showed the strongest algicidal effect and thus was selected for identification and further investigation of its effect on diatom nuclear DNA. On the basis of its morphologic and biochemical characteristics, it was classified as belonging to the genus Bacillus. The cells of this strain are rod-shaped \((0.75–1.3) \times (1.8–4.5) \mu m\), Gram-positive, immobile, and form elliptic spores (Figure 1A, Table 1). Microphotographs of ultrathin sections revealed the inclusions, ribosomes, and electron-transparent areas within the cells, as well as long curving or circular structures in the cytoplasm and near septae forming in the dividing cells (Figure 1B–G). Strain BS2-15 showed morphological and biochemical similarity to B. mycoides and B. weihenstephanensis (Figure 2). Comparative analysis of the nucleotide sequences demonstrated that it is closest to a strain of B. mycoides (acc. no AB021192), with 99% similarity. On the basis of the phylogenetic analysis (Figure 2) and main characteristics of the species (Table 1), this strain was named Bacillus mycoides BS2-15.
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**Figure 1.** Cells of *B. mycoides* BS2-15. (A) fluorescent microscopy; (B–G) transmission electron microscopy: (B) longitudinal section of a *B. mycoides* cell; (C) cross-section of a *B. mycoides* cell; (D) a fragment of Figure 1B showing the long curving structure in the cytoplasm; (E) a longitudinal section of a dividing cell; (F) a fragment of Figure 1E, septae are marked by the arrows; (G) a fragment of Figure 1E; black arrow marks a circular structure in the cytoplasm, white arrow marks the cell wall. Scale bars: (A) 5 μm; (B) 500 nm; (C,F,G) 200 nm; (D) 100 nm; (E) 1 μm.
Table 1. Morphological and biochemical features of strain BS2-15.

| Features              | BS2-15          | B. mycoides |
|-----------------------|-----------------|-------------|
| Motility              | −               | −           |
| Cell morphology       | rod-shaped      | rod-shaped  |
| Cell diameter, µm     | 0.75–1.3        | >1 µm       |
| Cell length, µm       | 1.8–4.5         | nd          |
| Gram status           | +               | +           |
| Endospores            | +               | +           |
| Elliptic              | +               | +           |
| Cylindrical           | −               | −           |
| Spherical             | −               | −           |

Enzymatic activity

|                | BS2-15 | B. mycoides |
|----------------|--------|-------------|
| Catalase       | +      | +           |
| Caseinase      | −      | +           |
| Gelatinase     | +      | +           |
| Lecithinase    | +      | +           |
| Lypase         | +      | nd          |
| Amylase        | −      | +           |

Acid production from carbohydrates:

|                | BS2-15 | B. mycoides |
|----------------|--------|-------------|
| Glucose        | +      | +           |
| Galactose      | −      | d           |
| Maltose        | +      | +           |
| Raffinose      | −      | −           |
| Lactose        | −      | −           |
| Sucrose        | +      | d           |
| Fructose       | +      | d           |
| Rhamnose       | −      | −           |
| Xylose         | −      | −           |
| Arabinose      | −      | −           |
| Dulcitol       | −      | −           |
| Inose          | −      | −           |
| Sorbitol       | −      | −           |
| Mannitol       | −      | −           |
| β-galactosidase| −      | d           |

—−this feature is absent; +−−this feature is present; d−−this feature is present in 80% of strains; nd—no data.

3.2. Algicidal Effect of B. mycoides BS2-15 on U. acus BK280

We demonstrated that different dilutions of the bacterial culture B. mycoides BS2-15 affect the diatom growth differently. A low density of added bacteria acted as a weak growth promoter of diatom culture (p = 0.01), whereas adding of bacteria at a high density (40–100 µL/mL) considerably decreased their growth (p < 0.01) (Figure 3).
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**Figure 3.** Inhibition of *U. acus* BK280 growth after inoculation with different concentrations of *B. mycoides* BS2-15 bacterial culture (microliters from optical density of bacterial culture of 0.9–1.0 at A600 nm).

The bacterial cell suspension added to *U. acus* is cultured in a 1% peptone water; therefore, it is critical to study its influence on diatom growth because data suggest that it can suppress their growth and result in their death [32]. As suggested in [32], such a reaction may occur because peptone is a mixture of dissolved free amino acids (DFAA). In a natural
environment, DFAA leakage can occur as a result of cell death and can hypothetically regulate the growth of diatoms. In this study, when peptone diluted to 100 µL/mL was added to *U. acus*, growth suppression was observed (*p* < 0.01) (Figure 4).

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*Figure 4.* Growth of *U. acus* BK280 during inoculation with 1% peptone solution (100 µL/mL), *B. mycoides* BS2-15 bacterial culture (100 µL/mL), bacterial filtrates (100 µL/mL) and bacterial pellet (bacterial culture washed from the 1% peptone solution, 100 µL/mL).

The assessment of the dynamics of cell mortality revealed that when culturing with 1% peptone solution, cell mortality increased compared with the control (*p* < 0.01); however, when culturing with bacterial culture, diatom cell mortality reached nearly 90% by the ninth day of culture and statistically significant differences were observed both with the axenic culture of diatoms and culture of diatoms cultivated with 1% peptone solution (*p* < 0.01) (Figure 5).

It is known that bacteria can manifest an algicidal effect either through physical contact with algal cells [33] or by producing algicidal compounds into the environment [15].

To determine the source of the algicidal effect, we added the filtrate of bacteria as well as bacteria washed from culturing medium (bacterial pellet) to the diatom cells (Figure 4). Bacteria added at a dilution of 100 µL per mL of diatom culture but washed from the medium did not affect the growth of the *U. acus*. Bacterial filtrate resulted in suppression of diatom growth (*p* < 0.01). It can be concluded that the algicidal substance is produced by bacteria in the environment, regardless of the presence of diatoms.

TEM has revealed that control cells had nuclei with large nucleoli, clearly visible envelopes, and nuclear pores (Figure 6A,B). The cross-sections of chloroplasts and mitochondria were similar to those observed previously (Figure 6A–C) [34]. Cells grown in the presence of *B. mycoides* also had normal chloroplasts and mitochondria similar to controls, but their nuclei often did not have clearly visible membranes on the side oriented to the cell center (Figure 6D,E). Even a series of slices from such cells did not contain clearly visible nuclear pores.
Figure 5. Mortality of the cells of *U. acus* BK280 after inoculation with 1% peptone solution and *B. mycoides* BS2-15 bacterial culture.

Figure 6. *U. acus* cells in axenic culture (A–C) and after culturing with *B. mycoides* BS2-15 bacterial culture (D–E) (TEM). (A,B) cross-sections of nuclei in different cells, with arrows marking nuclear pores; (C) a cross-section of a mitochondrion and a part of a chloroplast in a native cell; (D) a damaged nucleus of an *U. acus* cell under algicidal effect; (E) a cross-section of a mitochondrion and a part of a chloroplast under algicidal effect. Abbreviations are: Chl—chloroplast; m—mitochondrion; N—nucleus; Nc—nucleolus. Scale bar: (A,B,D) 500 nm; (C,E) 200 nm.
TUNEL-analysis demonstrated that in the axenic culture, nucleus fragmentation did not occur (Figure 7A). To model the DNA lesions (positive control), the cells were permeabilized with DNase I, which caused 90% of the nuclei to show DNA fragmentation (Figure 7B).

During co-cultivation of *U. acus* with *B. mycoides* BS2-15, numerous nuclei of diatom cells were already stained with Alexa Fluor™ 488 after 24 h, showing evidence of DNA fragmentation (Figure 7C). In addition, 5% of the nuclei appeared to contain both native and fragmented DNA. In such nuclei, separate regions were stained with the two dyes.

The cell valves of the axenic culture were intact (Figure 8A), as were the cell valves to which the bacterial filtrate was added (Figure 8D). When cultured with bacterial culture, many destroyed shells were encountered (Figure 8B), but also intact shells were preserved, but with *B. mycoides* cells attached to them (Figure 8C). It can be assumed that the bacteria consume the organic matter covering the valves of the diatoms, leading to increased fragility of the shells. Conversely, the algicidal effect is likely achieved by a complex of substances present in the environment and released by directly attached bacteria.
Figure 8. The state of a silicon frustules of *U. acus* under different conditions. (A) axenic culture; (B, C) co-culturing with *B. mycoides* BS2-15 bacterial culture; (D) adding of bacterial filtrate to diatoms.

It was previously demonstrated that a large proportion of the near-bottom layer of Lake Baikal is composed just by *U. acus* and diverse bacteria [9]. Because of the vertical water exchange phenomenon, live diatoms can precipitate at the bottom. It was also observed that bacteria from bottom sediments can use the cell wall of *U. acus* as a carbon source [35], and this corresponds to the observed increase of frustules fragility after co-culturing.

Stress conditions not only decrease the diatom growth rate but also result in lipid accumulation [36]. Axenic culture of diatoms contained whole chloroplasts and rare lipid droplets (Figure 9A). However, cells cultured with bacterial filtrate (Figure 9B) and bacterial cultures (Figure 9C) had no chloroplasts (indicating that the cells were dead) but contained numerous large and small lipid droplets.
It is known that nutrient deficiency in diatoms results in cell metabolism conversion to lipid accumulation [37]. We could suppose that diatoms cells compete with bacteria for nutrients, but the fact that lipids accumulated when diatoms were cultured with bacterial filtrate with the same result suggest that in this case, the mechanism of lipid accumulation is different. Also, the accumulation of elevated lipid results from the aging of the diatom culture and the effects of different chemicals [38,39].

4. Conclusions

*Bacillus mycoides* strain BS2-15 isolated from Lake Baikal bottom sediments produces a strong algicidal effect on the diatom algae *Ulnaria acus*. We revealed that algicidal effect does not depend on the presence of diatoms in a medium with bacteria, and bacterial pellets do not influence the growth of *U. acus*. We identified the fragmentation of the *U. acus* nucleus resulting from the co-culturing of bacteria and diatoms. Moreover, we demonstrated that DNA fragmentation at *U. acus* is observed after only 24 h of common culturing. The specific staining of these nuclei and decreased culture growth in the presence of *B. mycoides* BS2-15 suggests the presence of specific response mechanisms in diatom cells that are involved in the algicidal activity of the bacteria. This phenomenon highlights the complexity of algae-bacteria interactions, which can be involved in regulating microorganism populations in water ecosystems.

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