Abstract. Dendroflora is an important component of all urban ecosystems, acts as an accelerator of biogeochemical processes in the soil and serves as a filter to clean the air of toxic compounds. However, the progressive growth of the impact on the urban ecosystem invariably leads to a different kind of dendro-pathogenic problems caused by abiotic and biotic factors. Therefore, the aim of our study was the development of a biological product based on microorganisms – antagonists of phytopathogens and based on the results of microbiological and ecological studies to highlight indigenous strains of *Bacillus*, having antagonistic activity against a variety of phytopathogenic microorganisms, i.e. trunk rots pathogens of elm.

Keywords: dendroflora, ecosystem, antagonistic activity, enzymic activity, pathogenic, elm.

1. Introduction

Under present-day conditions, intensive growth of anthropogenic impact on urban ecosystem is a reason for the decline of core indicators of the human environment. Increase of the toxic compounds concentration in the air and in the soil reduces the vital signs of the trees, causing various diseases of physiological nature. On the other hand, strengthening of the process of uncontrolled importation of raw plant building materials and natural plant introductions contribute to invasion of new species of pathogenic organisms that cause deterioration of the phytosanitary status of woody communities in settlements. Dendroflora of South Kazakhstan province is an exclusive community consisting of more than 600 species and varieties of indigenous and introduced tree species. Existing forest stands in these communities are well adapted to the soil and climatic conditions of the south of Kazakhstan and play an invaluable role in regulating of settlement microclimate in arid climates. However, since the mid 90-ies of the 20th century there was started the trend of deterioration of dendro-pathogenic state of these communities: sick trees was increased in number in the region with previously unknown diseases that leads to complete loss of indigenous species of trees community. Elm was affected by the strongest degradation,
which is one of the main tree species. Currently, this fact is one of the major environmental problems in Southern Kazakhstan. Meanwhile, it is known that in conditions of urban ecosystem the chemical methods of plant protection are not applied, that justifies the need for the development of biological control methods against phytopathogenic objects.

_Ulmus parvifolia_ is one of the indigenous species in South of Kazakhstan. It contributes the large part of regional dendroflora. In recent years, phytosanitary condition of the regional dendroflora association has been sharply deteriorated especially as regards to population of _Ulmus pumilla_ L. This is connected with invasion of foreign pests-xylophage that are spreading spores of plant pathogenic fungus. Plant pathogenic fungus _Fusarium solani_ (Mart.) Sacc. is one of the pathogenic agent of this species in South of Kazakhstan that affects plant’s stems resulting in their damage (Alakonya, 2009). In the stem areas affected by pests and pathogens appears secondary microflora causing decay (Maiorano, 2009).

Under the conditions of urban ecosystem, use of biological products for spread of contagium in dendroflora appeared to be the most potential bearing method of control (Stenglein, 2009). Today, search of novel strains antagonists adopted to certain soil-climatic conditions is still to be a relevant study (Phae et al., 1990; Kim et al., 2006; Suga, 2008; Kosova, 2009; MacLeod, 2010). Therefore, in prior investigations we have isolated and identified 5 antagonistic strains of _Bacillus_ bacteria. Among them _Bacillus thuringiensis 4ant_ strain appeared to be the most active antagonist for several plant pathogenic fungus justifying its prospective potential to be used as an active agent in biological product. Therefore, the objective of our studies was to determine the level of antagonistic and enzymatic activity of _Bacillus thuringiensis 4ant_ strain.

In scientific literature it is known that one of the key factors for implementation of antagonistic potential is bacillus ability to produce antibiotics (Ongena et al., 2008). However, fungicidal activity have only some of them, e.g. plipastatin A and B, iturine, bacillomycine D, mycobacillin and several other (Solanki et al., 2012). In most cases, bacillus strains produce a number of antibiotics of similar structure while their synergistic action leads to suppression of wide range of microorganisms. In a number of papers marked higher performance of culture filtrate than cell suspensions (Thara & Gnanimanickham, 1994). Secretion of antibiotics is the main but far not the only one way for suppression of fungi formation and antagonism manifestation. Multiple studies of hydrolytic complex of bacillar enzymes have shown that main functional load in lysis of chitin of fungi cell walls is exercised by chitinase (Balhara et al., 2011).

Chitinase is produced by most of _Bacillus_ strains that are lytic to cell walls of plant pathogenic fungus. However, data available now does not allow to make a conclusion as regards to the role of chitinase in manifestation of antagonistic activity of bacillus. On the one hand, it was found a direct association between chitinase activity and ability to inhibit growth of micromycetes. On the other hand, in antagonistic activity of bacillus antibiotic compounds are prevailing while the role of chitinase complex being limited to utilization of chitin contained in soil (Berg, 2009; Wang et al., 2009; Preecha et al., 2010; Yuan et al., 2012). For example, in number of studies it is reported that among 1757 bacillus isolates only 12% was suppressing _Rhizoctonia solani_ J.G. Kühn, with 31% of them manifested chitinase activity (Alvares et al., 2011). The most probable is that bacillus fungicidal activity is determined by their simultaneous ability to produce and secrete to the ambient both antibiotics and mycolitic enzymes.

2. Objects and methods

An object of the study was _Bacillus thuringiensis 4ant_ strain isolated in soil samples taken from seven districts of South Kazakhstan region.

**Culture and fermentation.**

Bacteria was cultivated in the medium with the following composition: (g/l): potato starch 239, maize extract 12.43, yeast 2.19, lactose 2.87, sulphide of copper 0.0036, ammonium phosphate 8.4, sodium chloride 0.28, magnesium sulfate 0.14, potassium sulfate 2.87, calcium chloride 0.45, pH 6.2-7.0. Bacterial cells were cultured in 250 ml flasks with 40 ml of medium for inoculation and cultivated with orbital rotator 250 rpm during 12 hours at 28°C. Pure culture of phytopathogenic fungi was incubated in Chapek’s medium.

Level of antagonistic activity of bacteria: antagonistic activity of _Bacillus thuringiensis 4ant_ was revealed with volume displacement method (Egorov, 1994.) Assessment of antagonistic activity was performed at day 7 of incubation from the diameter of sterile areas in lawn fungus formed around the cavities. Enzymatic activity of bacteria: growth rate of bacteria in liquid medium was judged from variation of optical density detected with spectrophotometer SF-46 at λ=590 nm.

Total chitinase activity was defined from the amount of reductive sugar resulted from chitin hydrolysis with DNS reagent (DNS method) (Aktuganov, 2000). Specific activity was calculated as the ratio of the number of total enzymic activity in the culture fluid to optical density value of growth and expressed in relative units.

Phosphomonesterase activity was determined with e-nitrophenylphosphate (Leschinskaya, 1980) as a substrate. Activity was expressed in μM of broken substrate.
hydrolysable in enzyme solution 1 ml within 1 minute of incubation. For conversion was used calibration curve of relation of absorption of solution at $A=410$ nm with concentration of p-nitrophenol.

Nitrogenase activity of *Bacillus thuringiensis 4ant* was determined by acetylene method that is based on nitrogenase ability to reconstitute acetylene up to ethylene in amount proportional to the nitrogen with gas chromatograph (Kalininskaya et.al., 1989). Glucose autolysate medium was dispensed in 3.5 ml vials and inoculated by bacterial suspension. Seed stock was a rinse of daily culture from slope agar that was added to the medium up to the final concentration of 0.05 ml. Following a certain period of incubation at 28-30°C was performed neutralization with 0.1 N NaOH sterile solution of those vials where bromothymol blue discolored to yellow; cotton swabs were replaced with rubber stoppers clamped with metal clips. In parallel with control samples were prepared test samples (in 3.5 ml vials with glucose autolysate medium added distilled water equivalent to quantity of seed stock in test vials). Nitrogenase activity was expressed in $\mu$g N$_2$/ml*h. In determination of nitrogenase activity level the following medium and solutions were used:

1) Glucose autolysate medium, g/l: glucose – 10.0; yeast autolysate – 0.08; $K_2$HPO$_4$ – 1.74; KH$_2$PO$_4$ – 0.91; MgSO$_4$$\times$7H$_2$O – 0.3; NaCl 0.5; CaCl$_2$$\times$6H$_2$O – 0.1; FeCl$_3$$\times$6H$_2$O – 0.01.
2) Micronutrient mixture by Fyodorov, g/l: $H_2$BO$_3$ – 5.0; $Na_2$MoO$_4$$\times$2H$_2$O – 5.0; MnSO$_4$$\times$H$_2$O – 3.0; KJ – 0.5; NaBr – 0.5; ZnSO$_4$$\times$7H$_2$O 0.2; $Al_2$(SO$_4$)$_3$$\times$12H$_2$O – 0.3; bromothymol blue – 0.01-0.02.

3) Vitamins, $\mu$g/l: biotinum – 10.0; $B_12$ – 2.0; riboflavin – 200.0; thiamine, pyridoxine, calcium pantothenate, nicotinic and e-aminobenzoic acids – 100.0 each.
4) Solutions: 0.1 N NaOH.

3. Results and discussion

For the purpose of determination of antagonistic activity level of *B. thuringiensis 4ant* 9 archival strains of phytopathogenic microorganisms were used. Findings have shown that *B. thuringiensis 4ant* as regards to test-objects yielded highest antagonistic potential both in relation to archival and regional pathogenic agent of plants (Table 1).

In the Table 1 strain-antagonist exhibits its highest antagonistic activity against *F. solani, F. oxysporum, F. sambucinum* and *M. circinelloides*. These micromycetes are main pathogens of Ulmus parvifolia.

3.1. Enzymic activity of *B. thuringiensis 4ant*

Currently an intensive research of hydrolytic complex of bacillary enzymes (chitinase, protease, cellulase, glucanase etc.) (Liu, 2011) is conducted where chitinase plays main role in lysis of chitin of fungi cell walls (Velbo et al., 2011). We found that chitinase specific activity varied up to 22.8 relative units (Fig. 1).

Table 1. Antagonistic reaction indicators of *B. thuringiensis 4ant* strain against a number of phytopathogenic microfungus

| Phytopathogenic microorganisms | Zone of inhibition, mm |
|-------------------------------|-----------------------|
| *Fusarium solani* (Mart.)Sacc. | 24.0 ± 0.4            |
| *Fusarium oxysporum* Schlecht. emend. Snyder & Hansen. | 28.0 ± 0.2            |
| *Fusarium graminearum* (Schwein.) Petch, | 19.0 ± 0.6            |
| *Rhizoctonia solani* J.G. Kühn | 17.0 ± 0.3            |
| *Bipolaris sorokiniana* (Sacc.) Shoemaker | 16.0 ± 0.4            |
| *Fusarium sambucinum* Fückel, | 22.0 ± 0.4            |
| *Alternaria solani* Soraure | 16.0 ± 0.5            |
| *Xanthomonas campestris* Dowson | 18.0 ± 0.4            |
| *Xanthomonas oryzae* Swings et al. | 15.0 ± 0.6            |
| *Mucor circinelloides* Tieghem. | 23.0 ± 0.4            |
It is interesting to note that \textit{B. thuringiensis 4ant} exhibits chitinase activity and ability to suppress a wide range of phytopathogenic micromycetes. It gives grounds to suggest that manifestation of Bacillus antagonism may be relied upon secretion of complex of chitinolytic enzymes.

3.2. Phosphomonoesterase activity of \textit{B. thuringiensis 4ant} strain

\textit{Bacillus} strains can be included into the development of biological products being potential bearing microorganisms that in addition to manifestation of antagonistic activity can transfer problematic organic and inorganic phosphorus compounds into available forms for plants. Figure 2 shows variability of phosphatase activity at 6 hours of incubation – 0.551, 12 h – 0.303 and 18 h – 0.257 relative units.

Nitrogenase activity of \textit{B. thuringiensis 4ant}. For many of natural ecosystems with nitrogen deficiency microbiological nitrogen bonding is the only ecologically clean way of entering of nitrogen compounds that are available for plants (Wang et al., 2009). Findings have shown that nitrogenase activity in culture fluid of \textit{B. thuringiensis 4ant} at 8 and 16 hours of incubation to be $4.1 \times 10^{-2}$ and $5.9 \times 10^{-2}$ μg N/ml*h respectively (Fig. 3). Peak activity was recorded in the culture at 24 hours of growth ($12.0 \times 10^{-2}$ μg N/ml*h). Gradual decrease of the enzyme activity occurred with the more time of cultivation and at 96 h it was $3.4 \times 10^{-2}$ μg N/ml*h.
4. Conclusion

*B. thuringiensis 4ant* strain isolated from soil samples appeared to be an efficient antagonist of pathogen of stem disease *Ulmus pumila – F solani*. Additionally, it was found that this bacterial strain can inhibit growth of the number of micromycete-phytopathogenes such as *F. oxysporum, M. circinelloides, F. sambucinum*. Wide range of antagonist properties of this strain justifies its practical value for to be a useful agent in biocontrol of dendroflora diseases in South of Kazakhstan.

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