Entomopathogenic activity of bacterial and viral strains from the bioresource collection "State Collection of Entomoacariphages and Microorganisms"

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Abstract. As a result of a series of experiments to study the entomopathogenic activity of bacteria and viruses from the "State Collection of Entomoacariphages and Microorganisms", it was revealed that the large wax moth Galleria mellonella L. it is sensitive to new strains of bacteria and viruses. In the future, the strains can be used in the subsequent stages of screening of microorganisms that are promising as agents of biological pest control. The maximum biological efficacy on the fifth day against the tested insect was observed when using bacterial strains Bacillus spp. BZR 1159 (94.6%) and BZR 936 (95.0%) and a granulovirus strain of the codling moth (CpGV) BZR L-5 (100%). Larvae of G. mellonella L were susceptible to melanization in the process of infection with new bacterial and viral agents.

1 Introduction

Recently, more and more attention in Russia has been paid to biological plant protection. In this regard, the search for environmentally friendly methods for regulating the number of harmful organisms is of great theoretical and practical importance. One of them is the development of microbiological preparations based on entomopathogenic microorganisms [1-2]. Bacillus thuringiensis, B. popilliae, Lysinibacillus sphaericus, Paenibacillus larva, etc.) and viruses of nuclear polyhedrosis and granulosis are actively used as active substances [3-7]. The advantage of viral and bacterial insecticides is their environmental friendliness and cumulative effect in agrocenoses [8-9]. It should be noted that viral agents have specific toxicological activity against target insects, thus having a narrowly targeted effect only on the target insect [10-13].

The use of chemical insecticides, in addition to the destruction of harmful insects, is known to inevitably reduce the number of useful entomofauna, which leads to a violation of the ecological balance of agrocenoses. In addition, with the intensive use of chemical pesticides, phytophages have a risk of developing resistance [14, 15]. Bioinsecticides solve this problem by acting selectively and destroying a certain range of pests while maintaining

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natural balance. However, the range of biological insecticides existing today is rather narrow in comparison with chemical ones and requires expansion [16]. In this regard, the purpose of our research was to study the insecticidal properties of entomopathogenic bacteria and viruses, which in the future may become the basis of biopreparations against agricultural pests.

2 Materials and methods

The objects of research are aboriginal, first isolated from farming ecosystem of southern Russia, entomopathogenic strains of bacteria and viruses from the bioresource collection (BRC) of the Federal State Budgetary Scientific Institution "Federal Research Center of Biological Plant Protection" (FSBSI FRCBPP) "State collection of entomoacariphages and microorganisms."

The bacterial strains *Bacillus* spp. BZR G2, BZR 936, BZR 1159 were selected for their ability to synthesize a complex of hydrolytic enzymes (lipase, chitinase, and protease) and was assessed by the insecticidal activity *in vivo*. The granulovirus strains of the coding moth *Cydia pomonella* L. (CpGV) BZR 6, BZR 10, BZR L-5, BZR L-7 were selected according to the results of the previously conducted primary screening for the large wax moth [17].

Caterpillars of the laboratory population of the large wax moth *Galleria mellonella* L., grown in the laboratory of chemical communication and mass breeding of insects of the FSBSI FRCBPP, were used as a test object for studying the effectiveness of bacterial and viral cultures of microorganisms. Due to its physiological characteristics, it is mass-reared for scientific and practical purposes as a model object for research in terms of assessing the effectiveness of biopreparations, as well as a food base for predatory bugs, tachin flies, trichograms, etc. [18, 19].

Bacterial strains were plated in Petri dishes (PD) by streaking on meat-peptone agar (MPA). The cultivation was carried out in a thermostat at a temperature of +28 °C for 48 h. At the end of the cultivation, an aqueous suspension was obtained by the washout method, and the number of bacterial cells was determined by the deep method [21].

Aqueous suspensions of viruses were prepared from the affected caterpillars of the coding moth, pre-selected due to the presence of obvious signs of viroses (inactivity, lethargy, discoloration of the cuticle, softening or partial necrosis of tissues). The insects were placed in test tubes with 10 ml of sterile water per sample and left under room conditions for decomposition and natural release of the virus from the cells. Then the sample was crushed with a glass rod, filtered, and then used as a material for infection [22, 23]. To determine the titer of viral granules in the samples, the suspension was centrifuged on a 70% glycerol pad - 30 min., 8000 rpm. [23]. Then the samples were applied to a glass slide, fixed in a burner flame, and stained by Shvetsova method [24]. A Goryaev chamber was used to count the number of granules.

*G. mellonella* L. was bred in a thermostat at a temperature of 25-27°C, a relative humidity of 60-80%. Caterpillars were fed with a modified artificial nutrient medium [20]. Caterpillars of the 3-5 age were used for the laboratory tests.

Testing the entomopathogenic activity of bacterial strains *in vitro* was carried out according to the guidelines by Burov [25]. Caterpillars of large wax moth actively moving and showing no signs of disease. Healthy, actively moving individuals were selected for the experiment. In one repetition, 50 caterpillars and 25 g of diet were used, treated with 25 ml of the solution, which varied depending on the variant: in the control variant – water, in the experimental variants – suspensions of strains. The biopreparation Fermovirin YAP, SP (CpGV) and the biopreparation Lepidocide, SP (*B. thuringiensis* var. *kurstaki*, spore-crystal complex) were used as biological reference preparation. The experience is repeated three
times. During the test, the insects were incubated at a temperature of + 25-27 °C and a relative humidity of 60-80%.

The calculation of the biological efficacy was carried out according to the Henderson and Tilton formula [9], which takes into account the changes in the number in the experimental and control variants:

\[ E=100 \times \left(1 - \frac{OaKb}{ObKa}\right) \]  

(1)

where \( E \) – efficiency expressed as percentage of pest reduction adjusted for control; \( Ob \) – the number of live individuals before the treatment in the experiment; \( Oa \) – the number of live individuals after the treatment in the experiment; \( Kb \) – the number of live individuals in the control in the preliminary count; \( Ka \) – number of live individuals in control in subsequent counts.

Melanization of insects was assessed visually, by examining the hypodermis on the 1\textsuperscript{st} - 5\textsuperscript{th} days.

3 Results and discussion

The minimum insecticidal effect was observed in the Bacillus sp. BZR G2 strain on the 3\textsuperscript{rd} - 5\textsuperscript{th} days. Strains Bacillus spp. BZR 1159 and BZR 936 showed the maximum biological efficacy on the 5\textsuperscript{th} day: 94.6 and 95.0\%, respectively (Table 1).

| Option                        | Titre, CFU / ml | Biological efficacy, % |
|-------------------------------|----------------|------------------------|
|                               |                | 3\textsuperscript{rd} day | 5\textsuperscript{th} day |
| Bacillus sp. BZR G2           | (2.9 ± 0.3) x 10\textsuperscript{7} | 4.0 | 6.1 |
| Bacillus sp. BZR 936          | (2.2 ± 0.1) x 10\textsuperscript{8} | 89.3 | 94.6 |
| Bacillus sp. BZR 1159         | (6.0 ± 0.1) x 10\textsuperscript{9} | 93.3 | 95.0 |
| Lepidocide, SP (B. thuringiensis var. kurstaki, spore-crystal complex) | (1.9 ±0.1) x 10\textsuperscript{9} | 70.0 | 87.5 |

Thus, the insecticidal activity of new bacterial strains against the large wax moth G. mellonella L was confirmed.

Also, to assess the effect of bacterial insecticidal agents, signs of caterpillar melanization were evaluated in comparison with controls. Figure 1 shows the phased of the skin melanization comparing to the control individuals (no melanism, normal color).

In the control (Fig. 1-a), the hypodermis of the larva is of a typical cream color, caterpillars ranging in size from 2 to 2.5 cm. Melanization of the affected caterpillars began with characteristic black spots on cream-colored larvae (Figure 1-b). All dead caterpillars were melanized (black larvae, Figure 1-d).
In the course of the research, the sensitivity of the laboratory population of the large wax moth *G. mellonella* L. to new viral strains was assessed. The test results are presented in Table 2.

**Table 2.** Insecticidal activity and titer of aqueous suspensions granulovirus of the codling moth (*CpGV*) strains and a commercial viral preparation against larvae of the large wax moth *G. mellonella* L.

| Option                | Titre, granules / ml | Biological efficacy, % |
|-----------------------|----------------------|------------------------|
|                       |                      | 3rd day | 5th day |
| CpGV BZR 6            | (3.7 ± 0.1) x 10^6   | 2.0      | 20.0    |
| CpGV BZR 10           | (3.6 ± 0.3) x 10^6   | 27.0     | 60.0    |
| CpGV BZR L-5          | (7.7 ± 0.4) x 10^6   | 87.0     | 100.0   |
| CpGV BZR L-7          | (2.6 ± 0.2) x 10^6   | 52.0     | 58.0    |
| Fermovirin YAP, SP (CpGV) | (2.0 ± 1.0) x 10^9 | 0        | 11.0    |

As a result of assessing the pathogenicity of new viral strains, it was found that the maximum biological efficacy was observed in *CpGV* BZR L-5 strain and on the 5th day it was 100%. In other strains, insecticidal activity was lower. Minimal biological efficacy was observed in the variant treated with Fermovirin YAP, SP. Thus, the results of the experiment proved the insecticidal activity of new strains of viruses against the large wax moth *G. mellonella* L.

Melanization was also observed when exposed to viral strains and similar to the symptoms manifested during infection with bacteria. However, most insects were not completely melanized, most of them showed spots on the surface of the hypodermis, as in the case of infection with bacterial strains in the early stages. There was no significant visual difference between the studied strains and the reference preparation. Control insects remained the typical cream color throughout the research.

Thus, as a result of a series of experiments, it was revealed that the large wax moth *G. mellonella* L. is susceptible to new bacterial and viral strains with insecticidal activity, which in the future may become the basis of biopreparation against agricultural pests.
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