The Influence of Pesticide Carbocide on Soil Microorganisms

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Abstract—Pesticides enter the nature as a result of human economic activity and represent a major hazard for people’s health. As a result of studying production and experimental carbocide pesticide doses on soil microorganisms we have revealed the biodegradation capability of carbocide. In particular, it has been established that fungicide carbocide is exposed to microbial decomposition by bacteria species of Bacillus and Pseudomonas.

Keywords—pesticide, carbocide bacteria, soil microorganisms, decomposition, agent.

I. INTRODUCTION

One of negative issues of the present time is global chemical pollution of the biosphere that generates reasonable concern on possible ecological disruption in separate ecosystems [1, 2, 7]. The special danger is constituted by the synthetic nonnatural substances coming to the nature as a result of economic activity of a human [3]. The important place among them is occupied by the chemical crop and animal protection agents – pesticides, one of which is fungicide carbocide. Carbocide is the agent intended to protect from fungal diseases of vegetables, fruit and berry crops, and vineyards. It is recommended to protect vegetables from storage decay. The active fraction of agent is cuprum(II) tricaptolactan, bichloride, monohydrate. The generalized formula of the substance is C18H35Cl2CuN3O4. The molecular weight is 492.0. It is a green crystal substance with the melting temperature of 80–82 °C. It is very soluble in water, a xylene, chloroform, acetone and ethyl hydroxide. The maximum allowable concentration (MAC) in the air of the working area is 2 mg/m³. Working concentration of 1 mg of r.a. on 1 kg of soil. It is allowed for protection from mal secco, anthracnose, scab and gray mold (consumption rate of 9-12 kg of r.a./ha), to decrease losses at storage (including from gray rot) of sugar beet (10 g/t of agents [8]).

II. METHODS AND MATERIALS

In experiments there were used:

1) Samples of the black soil from fields of the farm enterprise “Bolatbi” in Kurchaloyevskyi district of the Chechen republic.

2) Fungicide carbocide - chlorine-containing heterocyclic hydrocarbon.
Advances in Engineering Research, volume 151

3) The cultures of microorganisms separated from the soil with pesticide: 7 strains of actinomycetes; 9 strains of heterotrophic bacteria; 5 strains of mold fungus [5].


There was used the scheme of methods in the study: soil sampling, preparation of the soil and pesticide to the research; entering the pesticide into the soil; sampling of soils and preparation of them to the research; defining dynamics of microorganisms population in the soil with pesticide (experiment) and without pesticide (control); studying the influence of pesticide on the “respiration” intensity of the soil, nitrogenic activity and nitrogen fixation; allocation of microorganisms strains from soil, containing pesticide for long period (30 days); determining sensitivity of pure growths to pesticide; selecting strains growing in the area with pesticide; studying of destructive activity of microbes.

Growing mediums for allocation and cultivation of microorganisms: M1A - meat infusion agar; SAA - starch-ammonia agar; WA - wort agar; Sabouraud environment; M9 - mineral carbon-free environment.

III. RESULTS

We have studied the influence of the working (1 mg/kg) and experimental doses of cartocide (10 mg/kg and 100 mg/kg) on changes of quantitative indices of microorganisms in the soil [4, 10].

The obtained results demonstrate that all doses of cartocide in the first days of the experiment stimulated heterotrophic bacteria, increasing their quantity by 1.5-2.5 times, in comparison with control (Fig. 1). Moreover, the stimulating effect was intensified by increasing the concentration of the agent. It means that the agent is undergone the degradation by soil microorganisms [6]. However, on the 30th day we revealed that the quantitative indices of bacteria had been decreased to control level. Perhaps, the cartocide was quickly available substratum for them, and it was abundantly used as nutritious and energy material. At the same time actinomycetes practically did not take up the agent. Only when a small dose of pesticide was introduced, the number of actinomycetes remained at the initial level. When introducing increased doses the inhibition of growing processes was observed, but on the 30th day their number was restored to control indices (Fig. 2).

Mold fungi were very sensitive to pesticide. We observed the suppression of their growth which lowered their quantitative indices to 70% (Fig. 3).

By 30 days in the experiment samplings containing working concentration of agent and containing 10 doses, there was weak renewal of mold fungus growth. However, the number of them nevertheless had not reached the control level yet.

![Heterotrophs](image1)

**Fig. 1.** The influence of different cartocide doses on the microorganisms population.

![Actinomycetes](image2)

**Fig. 2.** The influence of different cartocide doses on the microorganisms population.
Advances in Engineering Research, volume 151

10 doses fluence of different cartocide do.

teria to use representatives of this

cide, a source of carbon environment with pesticide processes. For this purpose the strains allocated from the soil group of geterotrophic bacteria we were planted to mineral agarized environment M9 with combinations of carbon substrate. As a source of carbon we added pesticide (P) or glucose (G). According to the environment containing glucose (M9+G) we studied the ability of bacteria to grow in the synthetic environment M9, but in the environment containing agent (M9+ P) we revealed the strains using pesticide as a substratum. To confirm splitting of substratum we used the bromthymol blue indicator.

**TABLE I. SENSITIVITY OF PURE GROWTHS MICROORGANISMS TO CARTOCIDE (1 MG/L)**

<table>
<thead>
<tr>
<th>Strain</th>
<th>Colony diameter, mm</th>
<th>Inhibition coefficient, %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
<td>Experiment</td>
</tr>
<tr>
<td>Bacillus sp. 2</td>
<td>7.5 ± 0.5</td>
<td>6.5 ± 0.8</td>
</tr>
<tr>
<td>Bacillus sp. 5</td>
<td>57.5 ± 0.5</td>
<td>46.5 ± 0.5</td>
</tr>
<tr>
<td>Bacillus sp. 4</td>
<td>6.5 ± 0.7</td>
<td>3.0 ± 0.6</td>
</tr>
<tr>
<td>Proteus sp. 5</td>
<td>6.5 ± 0.4</td>
<td>6.5 ± 0.5</td>
</tr>
<tr>
<td>Proteus sp. 7</td>
<td>12.5 ±0.5</td>
<td>12.0 ±0.8</td>
</tr>
<tr>
<td>Pseudomonas sp. 4</td>
<td>15.5 ± 0.4</td>
<td>20.0 ± 0.6</td>
</tr>
<tr>
<td>Pseudomonas sp. 6</td>
<td>8.0 ± 0.5</td>
<td>9.0 ± 0.5</td>
</tr>
<tr>
<td>Pseudomonas sp. 11</td>
<td>14.5 ± 0.4</td>
<td>15.5 ±0.8</td>
</tr>
<tr>
<td>Sporosarcina sp. 1</td>
<td>33.0 ±0.7</td>
<td>6.5 ± 0.6</td>
</tr>
<tr>
<td>Sporosarcina sp. 8</td>
<td>17.0 ±0.4</td>
<td>16.5 ±0.8</td>
</tr>
<tr>
<td>Staphylococcus sp. 9</td>
<td>15.0 ±0.6</td>
<td>7.0 ± 0.5</td>
</tr>
</tbody>
</table>

**TABLE II. SENSITIVITY OF ERWINIA STRAINS TO CARTOCIDE (1 MG/L)**

<table>
<thead>
<tr>
<th>Bacterial species and strain</th>
<th>Colony diameter, mm</th>
<th>Inhibition coefficient, %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
<td>Experiment</td>
</tr>
<tr>
<td>Envidia amylavora 75</td>
<td>6.4 ± 0.5</td>
<td>5.3 ±0.4</td>
</tr>
<tr>
<td>E. carotovora var. citrullis MH</td>
<td>11.7 ±0.2</td>
<td>1.5 ±0.5</td>
</tr>
<tr>
<td>E. carotovora var. citrullis 215</td>
<td>9.6 ±0.2</td>
<td>1.9 ±0.6</td>
</tr>
<tr>
<td>E. carotovora var. citrullis 438</td>
<td>6.4 ±0.3</td>
<td>4.5 ±0.5</td>
</tr>
<tr>
<td>E. carotovora var. citrullis 9011</td>
<td>5.4 ± 0.4</td>
<td>2.4 ±0.4</td>
</tr>
</tbody>
</table>

**IV. CONCLUSION**

We have studied 9 strains of heterotrophic bacteria, 60% of which are cultures of Bacillus bacteria, 40% of gram negative rod-like bacteria and cocci, and 5 strains of mold fungi. The obtained data (Tabl. 1) have allowed to reveal 5 strains of the heterotrophic bacteria using pesticide as the only source of carbon and energy.
References


