RESEARCH ARTICLE

Identification and Biological Properties of the Pathogen of Soft Rot of Tomatoes in the Greenhouse

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Abstract:
Recently, in the greenhouses of Ukraine, a sharp increase in the infestation of tomato plants with soft (wet) rot has been registered.

Purpose:
To identify the pathogen of the soft rot, study its biological properties and develop practical recommendations for plant protection measures.

Methods:
The work was performed using classical microbiological and biochemical research methods. The API 20E (Bio Merieux) test system was used to study the physiological and biochemical properties of the bacteria. The study of bacterial fatty acids was carried out by mass-spectrometry.

Results:
Using a bacteriological analysis of tomato plants selected in greenhouses of the Kyiv region, for the first time, virulent strains from this crop were identified and assigned, according to their physiological, biochemical, morphological and cultural characteristics, to the *Pectobacterium carotovorum* species. The fatty acid profile of the total cell lipids of the isolated strains was distinguished by the presence of 1,13‒1,17%, of 3-hydroxytetradecanoic acid, which is typical for *P. carotovorum* subsp. *carotovorum*. It was found that the isolated strains were sensitive to oxychloride and copper hydroxide-based fungicides with 10,5 to 17,5 mm growth inhibition zones.

Conclusion:
The causative agent of soft (wet) rot of tomatoes in greenhouses is *P. carotovorum* subsp. *carotovorum*. Fungicides with oxychloride and copper hydroxide are recommended as the active agents to protect tomato plants from the pathogen.

Keywords: Soft (wet) rot of tomatoes, *Pectobacterium carotovorum* subsp. *carotovorum*, Biological properties, Controls, Pathogen, Plant protection.

1. INTRODUCTION

In the vegetable growing nowadays, a steady trend of growth of bacterial plant diseases affecting various types of vegetable crops, is taking place. Unlike micromycetes, these pathogens are hardly predictable and cause significant crop losses. Tomato plants are affected by multiple bacterial pathogens, the spread of which is facilitated by changeable natural and climatic conditions as well as by the structural and functional specifics of the plants [1]. Over recent years, climate change has facilitated the proliferation of bacterial diseases into temperate climate zones and an increase in their severity. If the average daily air temperature in the summer increases by 3-4°C, the prevalence of bacterioses doubles and the infestation of plants grows by 30-50% [2].

Another important natural climate factor for the spread of bacterial diseases is increasing the length of the growing seasons and the frost-free periods. Under such conditions, the increase of occurrence of bacterial diseases is due to both typical and common in Ukraine pathogens and to the pathogens that previously had not been spotted on the territory of our
country. In particular, there is an increased degree of damage to tomato plants by such species as Clavibacter michiganensis subsp. michiganensis, Pectobacterium carotovorum, Pseudomonas corrugata, P. syringae pv. tomato, Xanthomonas vesicatoria and Pantoea agglomerans [3].

Favourable agro-meteorological conditions of the growing season are reported to facilitate the spread of the pathogenic bacteria of the genera Pectobacterium, Dickeya [4 - 6], which cause the soft (wet) rot of tomatoes. The disease is particularly damaging for tomatoes cultivated in the greenhouses, where up to 67% of the plants can become affected.

The pathogen P. carotovorum is reported to affect a wide range of host plants, in particular, potatoes, cabbage, lettuce, onions and carrots. It is widespread in various climatic zones, including the whole of Europe [7 - 12]. Most often, the lesion develops through small damage of the fruit skin or in the places where pests feed. When seedlings get damaged, the bacteria, under favourable conditions, can destroy the seedlings rather quickly. They penetrate the plant mainly through surface wounds; as a result, the stems get rotted most often near the substrate, and the fruits – near the pedicels. At first, a transparent spot appears on the affected fruit, then the spot gets pressed in and the skin cracks. The affected tissue of stems and fruits softens, liquefies and becomes dark brown; after 2–3 days, the fruit rots into a liquid mass with an unpleasant smell [13]. Similar symptoms are caused by the blackleg pathogens Dickeya dianthicola and D. solani, and such similarity significantly complicates their identification and differentiation. If the fruit is infected while it is on the plant, it gradually dries off, and ultimately, only the skin remains. When the fruit is open, the formation of necrosis of the seed shell and fruit vessels can be observed; this wet rot is similar to the necrosis of the core of the tomato stem, the pathogen of which is Pseudomonas corrugate [14].

Usually, the development of the soft rot starts with the secretion by phytopathogens of a set of exoenzymes, primarily pectinases and cellulases, which are known to depolymerize the cell wall of the plant. The infections caused by Pectobacterium lead to maceration of parenchyma cells through the secretion of pectate lyases enzymes, polygalacturonases, pectin lyases, pectin methylesterases, cellulases and proteases. Such maceration is followed by the development of symptoms of softening of the affected plant tissues. The maceration depends on the external temperature and humidity. Optimal temperatures for the development of the disease are 25 - 30 °C [15].

The lack of timely diagnostics and express test systems for identifying phytopathogenic bacteria in seeds, seed material and plantings, the use of non-traditional farming systems that do not control plant bacteriose, lack of modern, highly effective protective measures against fungi, but not bacteria and lack of plant protectants lead to a gradual accumulation of infected material in fields, seeds, and weeds [2]. Pectobacterium spp. is a taxon that consists of strains with a number of different phenotypic, biochemical and genetic indicators, with a wide range of host plants and pathogenicity systems, which makes it difficult to diagnose this pathogen and choose suitable methods and ways for protecting tomato plants. Currently, the control of the spread of soft rot pathogens is based on steaming or sterilizing the soil in the greenhouse, as well as reducing the proportion of nitrogen and increasing potash fertilizers, removing diseased plants in nurseries, reducing the temperature and relative humidity of the air, careful sorting and culling of tomato fruits damaged by diseases 3–5 days after the harvesting before storage [16]. Under industrial production conditions, fungicides are used against pathogens of bacterial diseases since there are no special preparations with antibacterial activity registered in Ukraine. For the control of the bacterial diseases of tomato plants, the preference is given to sulfite, oxchloride and copper hydroxide [17], which, however, have been only partially effective and have not ensured complete quality control of the soft rot pathogens. In this regard, the development of effective methods for detecting and identifying the pathogen is an important element in the overall plant protection and tomato seed certification programs.

The aim of this work is to identify, investigate the biological properties and develop practical recommendations for plant protection against the pathogen of soft (wet) rot of tomatoes.

2. MATERIALS AND METHODS

2.1. Plant Material

Plants with affected, liquefied and softened dark brown stems with a liquid mass and an unpleasant smell, and chlorotic or necrotic light yellow leaves with abundant watery spots were selected at commercial greenhouses in the Kyiv region.

2.2. Isolation of the Pathogen

For microbiological analysis, small pieces of affected tissue (capturing areas of affected and healthy tissue) were consistently washed with sterile running water and placed on potato agar in Petri dishes and incubated in a thermostat at 26–28 °C for 48–72 h. For comparative system studies, we used the type strain from the collection of the Department of Phytopathogenic Bacteria of the D. K. Zabolotny Institute of Microbiology and Virology at the National Academy of Sciences of Ukraine – P. carotovorum subsp. carotovorum (Jones 1901) Waldee 1945 emend. Gardan et al. 2003, UCM B-10757° (ATCC15713, ICMP 5702, NCPPB 312).

2.3. Morphological and Cultural Properties of the Pathogen

The morphology (size, shape, colour and growth patterns) of colonies was recorded after 24–48 hours of growth on a dish with a potato agar at +28°C. Gram staining was performed using a commercial IVD kit (Merck, Germany). The mobility of bacteria was determined by the “crushed drop” method on the one-day culture. The prepared slide was examined under binocular microscope Sigeta MB-201 (Sigeta, Ukraine) using immersion oil. Cell morphology was studied under a transmission electron microscope JEM-1400 (Jeol, Japan) at the Center for Open Access at the D. K. Zabolotny Institute of Microbiology and Virology at the National Academy of Sciences of Ukraine.
2.4. Kovac-oxidase Test

The commercial set of Oxidase Reagent (Bio Merieux, France) with N,N,N’,N’-tetramethyl-p-phenylene-diamine (TMPD) was used. Fresh bacterial culture (18–24 h) was spread over the TMPD-treated filter paper. Within 10-30 seconds, a dark blue or purple colour appeared, which confirmed the oxidation of the reagent.

2.5. Redox (o/f) Glucose Utilization Test (Hugh-Leifson Test)

A commercial set of Micro-La-Test (Erba Lachema) was used.

2.6. API Test System

To study the properties of the soft rot pathogen of tomatoes, API 20E test systems were used to identify gram-negative bacteria (BioMerieux, France). The results were estimated according to the results table, and identification was performed using the list of profiles API 20Е [18].

2.7. Pectolytic Activity Test

One-day cultures of separate isolates were transferred to sterile potato disks. The presence or absence of maceration was determined visually or by touching the disks with a bacteriological loop after one day of incubation.

2.8. Analysis of Fatty Acid (FA) Profile of Total Cellular Lipids

To obtain methyl esters, raw bacterial cells grown on the potato agar, were suspended in methanol with 1.5% of sulfuric acid. Methanolsis was performed in sealed ampoules at 80°C for 1 hour. The FA methyl esters were extracted with 3 ml of hexane. The extracts were evaporated on a vacuum rotary evaporator. Methyl esters of fatty acids were divided by Agilent 6890N/5973 inert mass-spectrometer system, HP-5MS column 30 m × 0.25 mm × 0.25 mm, temperature - 150-270°C with a gradient of 4 °C, carrier gas – helium, in the Open Access Center at the D. K. Zabolotny Institute of Microbiology and Virology of the National Academy of Sciences of Ukraine. The peaks were identified by comparing the time of their content with the time of the content of standard samples of FA methyl esters (Serva, Germany), as well as by the integrated mass spectrum database NIST 02. The FA content was determined as a percentage of the total peak area [19].

2.9. Study of Pathogenic Properties of the Pathogen

The ability to induce a hypersensitivity reaction was determined by injecting the suspension of bacterial cells with a titer of 10⁷ CFU/ml under the epidermis of tobacco leaves [19]. The pathogenicity of separated isolates was assessed by artificial infection of the stem and leaves of tomato plants [19].

2.10. Screening of Preparations with Antimicrobial Activity

The effect of the antimicrobial preparations on the tomatoes soft rot pathogen was studied by diffusion in a solidified potato agar. The experiment was performed in Petri dishes with holes made by cork drill in the center of the agar. The test preparations were added at the concentrations recommended by the manufacturer. The one-day bacterial suspension with a titer of 10⁷ CFU/ml was radially spread, and the dishes were incubated in a thermostat for 48 h at 28°C. The degree of sensitivity was determined by the diameters of the zones of test culture growth inhibition. If they were higher than 30 mm, the strains were susceptible, 15–30 mm – intermediate, if less than 15 mm – resistant [20].

Statistical calculations of the experimental data were performed using the Statistica 6.0 software.

3. RESULTS

The tomato plants picked from the greenhouse had blackening tissues in the basal aboveground part of the stem. On the cross-cutting section of the stem, the darkening of the vascular system and the damage of the parenchyma with rotting of the stem were observed. The leaves had specific wet, oily spots of various sizes. The fruits had watery gray-green or brown spots, and the plants had an unpleasant smell.

The bacterial samples were taken from the rotting stems of tomato plants on edge between the affected and healthy tissues (Fig. 1). Bacteria isolated from the affected plants formed colonies of white to cream colour on the potato agar. Most isolates did not cause maceration of potato disks, neither did they form dry rots. The gram-negative motile rods that could metabolise glucose under anaerobic conditions and cause maceration of potato pieces, specifically the appearance of brown watery spots at the site of application, were selected for subsequent research.

![Fig. (1). Tomato plant stems with soft rot lesions (natural infection).](image-url)
The pathogenicity tests showed that soft rot symptoms were developed within 72 h after the inoculation of tomato plants with the separated isolates (Fig. 2). It is worth noting that bacteria, re-isolated from artificially inoculated tomato plants, had the same morphological and cultural characteristics.

Since all the isolates virulent for tomatoes were similar in terms of morphological and cultural properties, we selected two isolates TKO-2 and TKO-7, for further identification. The TKO-2 and TKO-7 isolates, grown on the potato agar, formed translucent, shiny convex colonies of light gray colour with smooth edges 2–4 mm in diameter. An in-depth study of the biological properties of virulent isolates of bacteria showed that they were facultatively anaerobic single rods of 0,5–1,0 x 1–3 microns in size, motile by means of peritrichous flagella (Fig. 3), do not form spores, are oxidase-negative, catalase-positive, and have pectolytic activity.

The biochemical analysis of the TKO-2 and TKO-7 isolates has shown that growing on media with L-arabinose, D-glucose, D-sucrose and L-rhamnose formed an acid without gas. In addition to carbohydrates, the studied bacteria fermented β-galactosid and diluted gelatin (Table 1), which was similar to the typical strain of *Pectobacterium carotovorum* subsp. *carotovorum* UCM B-10757 and corresponded to the literature data for the bacteria of the species *P. carotovorum* [18]. Based on the results of the conducted biochemical analyses, the separated isolates were identified as the causative agent of the soft rot of tomatoes *P. carotovorum* subsp. *carotovorum*.

**Table 1. Physiological and biochemical properties of the causative agent of soft rot of tomatoes**

| Test/enzyme | Strains Isolated in Greenhouses in the Kyiv Region | *P. carotovorum* subsp. *carotovorum* UCM B-10757 | *P. carotovorum* according to the Literature Data |
|-------------|-----------------------------------------------|---------------------------------|---------------------------------|
| The form of cells | TKO-2, TKO-7 | Rods | Rods | Rods |
The study of the fatty acid composition of total cell lipids of the type P. carotovorum UCM B-1075 and selected strains of P. carotovorum subsp. carotovorum TKO-2 and TKO-7 confirmed the presence of fatty acids with carbon chain lengths C₁₂ - C₁₈, particularly: unsaturated – cis-9-hexadecenoic (C₁₆:1ω9c) and cis-11-octadecenoic (C₁₈:1ω9c), saturated – dodecanoic (C₁₂:0), pentadecanoic (C₁₅:0), hexadecanoic (C₁₆:0) and octadecanoic (C₁₈:0) (Table 2). Hydroxy acids were represented by 3-hydroxytetradecanoic acid (3-OH C₁₄:0) at a concentration of 1.13–1.17%, which is typical for soft rot pathogen P. carotovorum subsp. carotovorum.

For primary screening of antimicrobial activity against isolated bacteria, 15 chemicals were tested. In the experiments, preparations with the active substance copper hydroxide showed the greatest antibacterial activity against P. carotovorum subsp. carotovorum TKO-2 and TKO-7 strains, and growth inhibition zones ranged from 14 to 17 mm. Preparations with the active substance copper chloroxide showed moderate activity against P. carotovorum subsp. carotovorum TKO-2 and TKO-7 strains with growth inhibition zones from 10.5 to 13.5 mm. Most of the studied fungicides, recommended for processing plantations, did not affect the pathogen of soft rot of tomatoes (Table 3).

### Table 2. Fatty acid profile of total cell lipids of strains isolated from affected plants with signs of soft rot of tomatoes

| Fatty Acids             | Average amount of FA, % of the Total Content |
|-------------------------|---------------------------------------------|
|                         | TKO-2          | TKO-7          | P. carotovorum UCM B-1075² |
| **Formula**             | **Name**       | **TKO-2**      | **TKO-7**      | **B-1075²**    |
| C₁₂:0                   | dodecanoic     | 3.73           | 3.78           | 3.62           |
| 3-OH C₁₄:0              | 3-hydroxytetradecanoic | 1.17 | 1.13 | 1.14 |
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**Table 2** continued...

| Fatty Acids         | Average amount of FA, % of the Total Content |
|---------------------|-----------------------------------------------|
| C_{15:0} pentadecanoic | 1,77                                          |
| C_{16:1} cis-9-hexadecenoic | 33,40                                         |
| C_{16:0} hexadecanoic | 36,56                                         |
| C_{17:0} cyclo cis-9,10-methylenehexadecanoic | 1,37                                          |
| C_{18:1} cis-11-octadecenoic | 21,22                                         |
| C_{18:0} octadecanoic | 0,78                                          |

Table 3. Antibacterial activity of chemical protection against pathogens isolated from infected tomato plants in the Kyiv region

| Preparations of Fungicidal Action, Concentration | Diameter of Zone of Inhibition, (mm, M±SE) | Sensitivity Pattern |
|-------------------------------------------------|---------------------------------------------|---------------------|
| Copper chloroxide, 350 g/l                      | TKO-2: 10,5±0,4; TKO-7: 13,0±0,5             | Intermediate        |
| Kresoxim-methyl, 500 g/l                        | 0                                            | Resistant           |
| Aluminium tris (ethyl phosphonate) 75%, folpet 25% | 3,0±0,08; TKO-7: 3,5±0,08                     | Resistant           |
| Copper hydroxide, 770 g/l                       | TKO-2: 14,0±0,2; TKO-7: 17,0±0,5              | Intermediate        |
| Sulfur, 800 g/l                                | TKO-2: 3,0±0,08; TKO-7: 0,0                   | Resistant           |
| Metalaxyl, 80 g/l, mancozeb, 640 g/l           | TKO-2: 8,0±0,4; TKO-7: 6,5±0,2                | Resistant           |
| Azoxystrobin, 250 g/l                          | 0                                            | Resistant           |
| Difenconazole, 250 g/l                         | TKO-2: 7,5±0,3; TKO-7: 7,5±0,3                | Resistant           |
| Thiophanate-methyl, 500 g/l                    | TKO-2: 7,5±0,3; TKO-7: 7,5±0,2                | Resistant           |
| Mancozeb, 800 g/l                              | TKO-2: 7,5±0,2; TKO-7: 7,5±0,3                | Resistant           |
| Benomyl, 500 g/l                               | TKO-2: 3,0±0,08; TKO-7: 3,0±0,06              | Resistant           |
| Mancozeb, 302 g/l, propamocarb hydrochloride, 248 g/l | TKO-2: 6,0±0,2; TKO-7: 8,5±0,2                | Resistant           |
| Phenhexamide, 500 g/l                          | 0                                            | Resistant           |
| Thiamethoxam, 250 g/l, acetamiprid, 100 g/l, cymoxanil, 250 g/l, metalaxyl, 100 g/l | TKO-2: 7,5±0,2; TKO-7: 6,5±0,3 | Resistant |

Note: Resistant – < 15 mm; Intermediate – 15–30 mm; Susceptible – >30 mm

4. DISCUSSION

Pathogens of bacterial soft (wet) rot (SRP) belong to the genera *Pectobacterium* and *Dickeya*. These necrotrophic bacterial pathogens are spread widely amongst various farm crops. The pathogens can be isolated from infected plants, soil, and water; they can also be found on the surface and inside insects [21]. Due to the fact that SRP can significantly impair the growth, development and productivity of the plants, these bacteria have been attributed to the ten most harmful bacterial pathogens in agriculture [22]. For tomatoes under storage and transport conditions, the main pathogens of soft rot (rotting of fruits, tomato stalks, black leg) are *P. carotovorum* subsp. *carotovorum*, *P. atrosepticum* [21] and several species of *Dickeya* [23]. In Europe, blackleg and soft rot diseases can cause relatively high tomato crop losses both in the field and in the storage. Moreover, some studies have confirmed that *P. atrosepticum* and *P. carotovorum* subsp. *carotovorum* are among the most frequent pathogens causing soft rot and/or black leg in the temperate climate zones [6, 10, 24-26]. The symptoms caused by different SRPs in tomatoes are often hard to distinguish. SRP bacteria can cause systemic infection and death of the host plant. They usually produce a significant amount of extracellular enzymes that destroy the cell wall of the plant (for example, cellulases, pectinases, pectin methyl esterases, pectate lyases, polygalacturonases, phospholipases, proteases) and allow bacteria to enter the vascular compartments and lead to maceration (rotting) of the host plant tissues [22, 23]. Among all *Pectobacterium* spp. species that infect plants, *P. carotovorum* subsp. *carotovorum* has the widest range of hosts in the world, while *P. atrosepticum* is associated mainly with tomatoes grown in temperate climate [21]. Bacterial pathogens of soft rot cause watery lesions, rotting, softening of tissues and wilting of parts of the host plant, in particular stems, leaves, fruits and tissues of tomatoes [27, 28]. The bacteria isolated from the affected tomato plants were found similar to the species *P. carotovorum* subsp. *carotovorum*. According to the identified morphological characteristics, the isolates were facultative anaerobes, gram-negative, straight mobile rods, these features were consistent with previous experiments [29]. These isolates showed a positive hypersensitivity reaction, but some authors reported that *P. carotovorum* strains caused negative and positive effects of the hypersensitivity test on tobacco [30]. The isolates revealed high pectolytic activity, which is essential in the pathogenesis of this genus [31].

The choice of effective methods of protection against...
bacterial diseases is complicated by the similarity of the symptoms and by the cross infestation of growing plants with several different pathogens. Rapid withering away from plant tissues, leaf spotting and rotting are the main manifestations of damage by various species of the genus Pectobacterium. In order to facilitate the identification and investigation of the properties of isolates P. carotovorum subsp. carotovorum TKO-2 and TKO-7, the API 20E test system (BioMérieux, France), was used. It is based on 21 standardized biochemical tests for identifying phytopathogenic facultatively anaerobic gram-negative bacteria. The results of analysis showed that isolates of P. carotovorum subsp. carotovorum had similar biochemical characteristics. Nine of the twelve generally accepted API 20E tests were negative, in particular, nitrate reduction, indole production, urease, acetoin, hydrogen sulfide, arginine dehydrolase, lysine decarboxylase, ornithine decarboxylase, tryptophan deaminase. Whereas the isolates showed the ability to hydrolyze β-glucosidase, gelatins and decarboxylase, tryptophan deaminase. The presence of 3-hydroxytetradecanoic acid was found in both strains, which is a characteristic feature of the family Pectobacteriaceae. The results obtained regarding the fatty acid composition of P. carotovorum cell lipids isolated from tomatoes affected by soft rot are consistent with the literature data. The mechanism of copper ions action involves the denaturation of proteins, destroying enzymes that are crucial for the functioning of bacterial cells [17]. The work by Pradnyarani et al. [36] showed high antibacterial activity of copper hydroxide on the growth of P. carotovorum subsp. carotovorum bacteria under in vitro conditions. Preparations based on copper sulfate were characterized by an average value, while carbendazim and mancozeb were characterized by the absence of zones of bacterial growth suppression in the tested concentrations.

Copper compounds, as before, remain indispensable agents in the fight against bacterial pathogens in vegetable growing, although the literature describes some of the risks of their constant use, which are associated with the accumulation and contamination of soils and a high content of their residues in food products. On the other hand, copper plays a special role as an important element in the physiological and biochemical processes of higher organisms [17]. Therefore, the use of copper in all its varieties must be controlled and correspond to the environmentally friendly conditions and needs of plants.

**CONCLUSION**

On the basis of morphological, cultural, physiological and biochemical characteristics, isolates TKO-2 and TKO-7 of phytopathogenic pectolytic bacteria from the Ukrainian greenhouse were attributed to the species P. carotovorum subsp. carotovorum. On the basis of the verification of the antibacterial effect of plant protection products, it is recommended to use preparations with active substances oxyzchloride and copper hydroxide to fight the development of the pathogen of soft rot of tomatoes – P. carotovorum subsp. carotovorum.

**ETHICS APPROVAL AND CONSENT TO PARTICIPATE**

Not applicable

**HUMAN AND ANIMAL RIGHTS**

No animals/humans were used for studies that are the basis of this research.

**CONSENT FOR PUBLICATION**

Not applicable.

**AVAILABILITY OF DATA AND MATERIALS**

Not applicable.

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CONFLICT OF INTEREST

The authors declare no conflict of interest, financial or otherwise.

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