Identification and characterization of *Pseudomonas syringae* pv. *mori* affecting white mulberry (*Morus alba*) in Poland

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Abstract The white mulberry (*Morus alba* L.) is particularly valued in Asia and Europe because its leaves serve as fodder for the mulberry silkworm (*Bombyx mori* L.) in the production of raw silk (sericulture). However, the health of silkworm caterpillars producing silk fibres, and the quality of their cocoons depend strongly on fodder quality. Moreover, this plant is also valuable in the food and pharmaceutical industries, where high-quality herbal material is required. In one of the Polish mulberry plantations, trees with chlorotic, angular leaf spots and blotches were observed. The disease multiplied in spring in humid conditions and spread to developing blossoms and leaves. Leaves were collected during two growing seasons (2017–2018) from 18-year-old trees cultivated at the Experimental Farm of the Institute of Natural Fibres and Medicinal Plants in Petkowo, Poland (52°12′40″N 17°15′31″E). Leaf lesions exhibited bacterial streaming, and bacterial colonies were readily isolated on tryptic soy agar. Biochemical (Biolog Gen III system v. 2.8.0), molecular (16S rDNA and *gyrB* genes), and pathogenicity testing in the greenhouse identified the causal agent as *Pseudomonas syringae*. This bacterial species was the most abundant in each of the sample tested. To our knowledge this is the first description and characterization of *P. syringae* affecting mulberry trees not only in Poland but throughout Europe. Our aim was to point out the importance of plant diseases caused by bacteria. This finding serves as a warning for the phytosanitary services in Europe and for the sericulture industry that a new disease threat has appeared for European mulberry crops.

Keywords *Pseudomonas syringae* · Mulberry · *Morus alba* · Plant pathogenic bacteria · First report · Bacterial blight

Introduction

The white mulberry (*Morus alba* L.) is particularly valued in Asia and Europe because its leaves serve as fodder for larvae of the mulberry silkworm (*Bombyx mori* L.) in the production of raw silk (Barbour et al. 2008). Sericulture is a very important sector of the silk industry, but the health of silkworm caterpillars producing silk fibres, and the quality of their cocoons, depend strongly on fodder quality (Chen and Lu 2018). Any impairment of mulberry leaves caused by either microbial pests or plant protection chemicals affects negatively on the resulting silk quality (Chairman et al. 2012).
Moreover, mulberry leaves are very valuable in the food and pharmaceutical industries, where high-quality herbal material is required.

Mulberry trees have no special soil requirements and endure drought stress quite well. Moreover, this species is affected by a low number of natural pest, which limits the use of plant protection products and supports organic farming. Among mulberry arthropod pests are sap-sucking species, defoliators and borers belonging to the Lepidoptera, Hemiptera, Coleoptera, Thysanoptera, Orthoptera, Isoptera and mites (Rahmathulla and Sathyanarayana 2015; Wulandari et al. 2018). However mulberry may be affected also by various phytopathogenic agents such as nematodes (e.g., Meloidogynine incognita) (Muthulakshmi et al. 2010), phytoplasmas (e.g., mulberry dwarf phytoplasma) (Dai et al. 1997), fungi causing leaf-spots (e.g., Cercospora spp. and Mycosphaerella mori) (Barbour et al. 2008), and bacteria (Lou et al. 2011; Sahin et al. 1999; Wang et al. 2008).

Among bacterial diseases, bacterial blight caused by Pseudomonas syringae is a common disease of mulberry that has been reported from various agroclimatic conditions in Australia, Iran, South Africa, America, Indonesia, India and Pakistan (Akhtar and Sarwar 1988; Baharuddin 1997; Gupta et al. 1995; Wulandari et al. 2018). Infection by P. syringae can cause death of seedlings, a significant problem in mulberry nurseries (Hinrichs-Berger 2004), and reduces fruit yield and quality of mature trees (Bultreys and Kaluzna 2010). In general, studies of Pseudomonas strains from fruit trees are hampered by the fact that a host often supports multiple species (Mansfield et al. 2012). The taxonomic relationships of such fruit tree strains have been assessed in several genetic studies, leading to the designation of new pathovars (Kennely et al. 2007). At present P. syringae has been divided into over 50 pathovars, which are pathogenic to over 180 plant species, including woody plants, crops and grasses (Kaluzna et al. 2010; Khezri and Mohammadi 2018).

In Europe, P. syringae on fruit trees such as some fruits, hazelnut or citrus has been reported from France, the United Kingdom, Germany, Lithuania and Poland, however all reports concerned stone fruit trees, hazelnut, and citrus blast (Kaluzna et al. 2010). Currently, Pseudomonas syringae affecting mulberry trees has not been recorded in Europe, although it was reported in other parts of the world as P. syringae pv. mori, the most important bacterial disease of mulberry, causing bacterial blight (Baharuddin 1997; Gupta et al. 1995; Sahin et al. 1999; Takahashi 1980).

In 2017 and 2018, in one of the Polish mulberry plantations, trees with leaf symptoms resembling bacterial blight were observed. The aim of this study was to isolate, identify, and characterize the causal agent of the white mulberry disease using biochemical, molecular, and pathogenicity testing.

Materials and methods

Plant material

Symptomatic leaves of the Polish white mulberry cultivar Zolwinska wielkolistna were collected during two growing seasons, 2017 and 2018. The leaves were taken from 18-year-old trees cultivated at the Experimental Farm of the Institute of Natural Fibres and Medicinal Plants in Petkowo (52°12′40″N 17°15′31″E), and sent to the laboratory for the investigation. Four samples were collected, each consisting of 10 leaves collected from trees displaying chlorotic, angular leaf spots and blotches.

Pathogen isolation

The leaves were washed in tap water and surface-disinfested by washing in sodium hypochlorite (10%) for 1 min, a second washing in ethyl alcohol (70%) for 1 min, followed by triple washing in sterile distilled water. Next the leaves were homogenized in sterile physiological saline (0.9% NaCl) and investigated in the following manner. First, before homogenization, the symptomatic tissue was investigated under a light microscope (40 X) to check for bacterial streaming. The second test was electron microscopy screening for the presence of virus particles (Golding et al. 2016). Third, an aliquot of the homogenized leaf tissue was plated on potato dextrose agar (PDA; Sigma Aldrich Ltd.) to check for the presence of fungi, and lastly, the homogenized tissue was plated on tryptic soy agar (TSA; Sigma Aldrich Ltd.) to grow potentially present plant pathogenic bacteria.

The incubation lasted for five days on TSA medium at 27 °C to account for potentially slow-growing plant pathogenic bacteria. After incubation, a series of purification streaks was performed until a pure culture of each grown morphotype was obtained. The purity of each tested colony was verified on the basis of colony morphology and with Gram staining under the light microscope.
Pathogen identification

Both biochemical and molecular methods were used to identify the bacteria. For biochemical identification, the Biolog’s Gen III system (database v. 2.8.0) (BIOLOG Inc. Hayward, CA) was used, according to the manufacturer’s instructions. The molecular identification was based on the analysis of partial sequences of 16S rDNA and gyrB genes. Bacterial genomic DNA was isolated using a CTAB protocol (García-González et al. 2018), and the extract was kept at −20 °C for further experiments. For PCR amplification, the following universal bacterial 16S rDNA primers were used: 16S04 (AACTCAAA GGAATTGACGG) (Gerischer 2008) and pH 2 (AAG GAG GTG ATC CAG CCG CA) (Edwards et al. 1989). For the gyrB gene partial nucleotide sequence amplification, the 1480F/2242R primer pair (Bonasera et al. 2014) was used. PCR products of ~1400 bp for 16S rDNA and ~900 bp for gyrB were obtained using a touch-down PCR protocol (Korbie and Mattick 2008). The expected size amplicons were extracted from the agarose gel using Wizard SV Gel and PCR Clean-Up System (Promega, Madison, WI), and sequenced (Genomed S.A., Warsaw, Poland). The obtained sequences were verified against the GenBank database (BLAST), aligned, and the consensus sequences were submitted to the GenBank database to obtain accession numbers. For obtained DNA sequences, multiple sequence alignments were performed using ClustalW software, and a phylogenetic tree was constructed in Mega X software using the maximum likelihood method (bootstrapping 1000 repeats) (Kumar et al. 2018). The tree was rooted with Pectobacterium carotovorum 16S rDNA (NR_025316.1) and gyrB (FJ652727.1) sequences. The computed dendrograms contain all accession numbers and the names of the strains used in the analysis.

To confirm our identification results, the previously designed and published Pseudomonas syringae species-specific PCR primer pair (D21/D22) was used, as described (Schaad et al. 2001). Obtained amplicons of the expected size were analysed, sequenced, and submitted to the GenBank database as described above for the 16S rDNA and gyrB sequences.

Pathogenicity tests

All bacterial isolates obtained were tested in a pathogenicity test in greenhouse conditions (temperature 20–25 °C, humidity 50–80%) to confirm Koch’s postulates. Each isolate was tested on five healthy mulberry seedlings of the Zolwinska wielkolistna cultivar. The plants were inoculated with an aqueous bacterial suspension (10^6–10^7 cfu/ml) applied under high pressure on both sides of the leaves with a compressor sprayer, after wounding the leaves by abrasion with carborundum (EPPO Bulletin 2006). After inoculation, each plant was kept in a humid chamber for 72 h, and then in the greenhouse until symptom development. After incubation, symptomatic plants were subjected to reisolation to confirm Koch’s postulates. All reisolated strains were investigated as described above. Sterile distilled water sprayed instead of the bacterial suspension was used as a negative control, and an aqueous suspension (10^6–10^7 cfu/ml) of Pseudomonas syringae pv. syringae 2238, from the Institute of Plant Protection Bank of Pathogens collection was used as a positive control.

Since strains of P. syringae are often present as epiphytes on leaves (Arnold and Preston 2019), before performing the pathogenicity tests the leaves of the healthy mulberry plants to be used were washed with sterile distilled water and the suspension was plated on King’s B medium (Sigma Aldrich Ltd.) to ensure that the concentration of epiphytic Pseudomonas was uniform among plants.

Results

Symptoms description

The original symptoms observed on mulberry trees involved light-chlorotic lesions spreading on the leaf surface (Fig. 1). With time, the lesions increased in size from between 2 and 3 mm to 8–10 mm, their margins turned brown, and the centre of the lesions turn necrotic.

Pathogen identification

Biochemical identification

No virus particles were observed by electron microscopy. Also no fungal growth was observed on PDA medium. Hence, the presence of viruses and fungi as potential disease causal agents was excluded. Further tests were directed towards detection and identification of bacteria.

The examination of fresh, symptomatic leaf tissue under the light microscope showed a slight streaming of bacterial ooze. This feature is characteristic for most of the plant bacterial pathogens.
In total, 22 isolates were obtained during the two growing seasons. In 2017, 12 isolates were obtained, and 10 isolates were obtained in 2018. The pure cultures of all 22 bacterial isolates were subjected to biochemical identification using Biolog’s Gen III system. As a result the following bacterial species were tentatively identified: *Pseudomonas syringae* (four isolates), *P. fluorescens* (1), *Pantoea agglomerans* (4), *Serratia liquefaciens* (1), *Rothia mucilaginosa* (1), *Curvibacterium luteum* (1), *Cellulomonas cellaceae* (2), *Bacillus ruris* (1), and seven isolates resulted as “no identification”, indicating that they are not included in Biolog’s database (v. 2.8.0) that contains 1568 bacterial species comprising 258 plant pathogenic bacteria, including *Pseudomonas syringae*. The most abundant species in each of the four leaf samples tested, was *P. syringae*. Moreover all mulberry plants that developed symptoms in the pathogenicity tests turned out to comprise this pathogen (Table 1).

Molecular identification

Molecular identification based on partial nucleotide sequences of the 16S rRNA and *gyr* B genes confirmed this identification result (Table 1). The detailed results of BLAST verification of the obtained pathogen sequences are presented in the supplement (Table S1 and Table S2). A sequence identity matrix analysis of obtained 16S rDNA and *gyrB* sequences was performed (Tables 2, 3), and dendrograms of those sequences were generated using the maximum likelihood method and involving nucleotide sequences of both genes of the four detected *P. syringae* strains (Ma001, Ma011, Ma013, and Ma020), the three *P. syringae* strains obtained from reisolation (Ma011–1, Ma020–1, and Ma020–3), and other *Pseudomonas* plant-pathogenic and non-pathogenic environmental species sequences derived from GenBank. The accession numbers of all analysed sequences are included in the dendrograms (Figs. 2, 3).

Additionally, pathogen identification was verified by using and sequencing the products of the PCR primer pair D21/D22, which was published as *P. syringae*-specific, (Schaad et al. 2001). All tested pathogen samples resulted in *P. syringae*, confirming the identification based on 16S rDNA and *gyrB* sequencing (Table 1 and Table S3).

The percentage DNA identity between the sequences we obtained from the Polish mulberry isolates and the GenBank derived sequences of 16S rRNA and *gyrB* genes of *P. amygdali* pv. *mori* (syn. *P. syringae* pv. *mori*) is high, 99,2–100%, for 16S rRNA (Table 2) and 99–99,6% for *gyrB* (Table 3). This suggests that we detected *P. syringae* pv. *mori*, and that this is the actual causal agent of the disease, because most often, when a new disease emerges, the pathogen is clonal and at least all *gyrB* sequences would be expected to be identical or almost identical to each other, which is what we observed (Table 2, 3, Figs. 2, 3).

Pathogenicity test

The symptomatic leaves collected from the field were homogenized and all obtained bacterial isolates were used for inoculation of mulberry seedlings in the pathogenicity test, as described above. All symptom-causing isolates
turned out to be *P. syringae* (Table 1). After artificial inoculation, disease symptoms were observed for four out of 22 isolates tested; in each tested leaf sample (M1–2/2017 and M1-M2/2018, Table 1), the most abundant isolate was *P. syringae*. The symptoms developed within 7–10 days, whereas the negative control showed no disease symptoms. Indeed, the inoculation method used, spraying the leaves with bacterial suspension of the four isolates under pressure, resulted in obtaining disease symptoms resembling those observed originally in the infected trees (Fig. 4). The results obtained in the pathogenicity test allow to conclude that *P. syringae* is the actual causal agent of the disease of mulberry trees in Poland.

The observed pathogenicity of Polish *P. syringae* strains was much higher than the control *P. syringae* pv. *syringae* 2238 strain (Fig. 4).
This observation supports the thesis that Polish strains belong to *P. syringae pv. mori*, and that this is the causal agent of the disease since it elicits more severe symptoms than the *P. syringae* subsp. *syringae* strain used as the positive control. We are aware that using a high enough concentration of *P. syringae* and high pressure spraying under conditions conducive to disease, many *P. syringae* strains can cause some disease symptoms on several plant species even if they cannot cause disease on that plant under field conditions. Therefore, the result observed for the positive and negative controls is very important and supports our thesis.

**Discussion**

*Pseudomonas syringae* is an example of a plant pathogenic bacterium with significant impact, because of its basic biology and its impact on the agricultural economy. *P. syringae* is considered the first of the top 10 most important plant pathogenic bacteria in plant pathology (Mansfield et al. 2012). This pathogen has an extremely wide host range. That is why it is still debatable if the term “pathovar” is best description of *P. syringae* or whether we are dealing with one extremely versatile and adaptive single bacterial species (Mansfield et al. 2012). Its economic importance has been studied for a century, such as in case of bacterial canker of stone fruits. The results of such studies are extremely important because they inform epidemiological studies and the role of epiphytic stages in the life cycles of plant pathogenic bacteria (Bultreys and Kaluzna 2010).

*P. syringae* affects mulberry trees. Indeed, bacterial blight of mulberry caused by *P. syringae pv. mori* has been recorded in India, Pakistan, South Africa, Australia, Indonesia, and Turkey so far (Akhtar and Sarwar

### Table 2

|       | Ma001       | Ma011       | Ma013       | Ma020       | Ma011–1     | Ma020–1     | Ma020–3     | AB001446   |
|-------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|------------|
| Ma001 | ID          | 0.996       | 0.994       | 0.994       | 0.992       | 0.992       | 0.992       | 0.992      |
| Ma011 | 0.996       | ID          | 0.998       | 0.998       | 0.996       | 0.996       | 0.996       | 0.996      |
| Ma013 | 0.994       | 0.998       | ID          | 1000        | 0.998       | 0.998       | 0.998       | 0.998      |
| Ma020 | 0.994       | 0.998       | ID          | 1000        | 0.998       | 0.998       | 0.998       | 0.998      |
| Ma011–1| 0.992      | 0.996       | 0.998       | 0.998       | ID          | 1000        | 1000        | 1000       |
| Ma020–1| 0.992      | 0.996       | 0.998       | 0.998       | 1000        | ID          | 1000        | 1000       |
| Ma020–3| 0.992      | 0.996       | 0.998       | 0.998       | 1000        | 1000        | ID          | 1000       |
| AB001446| 0.992 | 0.996       | 0.998       | 0.998       | 1000        | 1000        | 1000        | ID         |

**Table 3**

|       | Ma001       | Ma011       | Ma013       | Ma020       | Ma011–1     | Ma020–1     | Ma020–3     | AB016355   | AB016347   |
|-------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|
| Ma001 | ID          | 0.994       | 0.978       | 0.991       | 0.993       | 0.996       | 0.996       | 0.993       | 0.990       |
| Ma011 | 0.994       | ID          | 0.983       | 0.993       | 0.994       | 0.998       | 0.998       | 0.995       | 0.991       |
| Ma013 | 0.978       | 0.983       | ID          | 0.980       | 0.978       | 0.981       | 0.981       | 0.985       | 0.981       |
| Ma020 | 0.991       | 0.993       | 0.980       | ID          | 0.993       | 0.994       | 0.994       | 0.991       | 0.988       |
| Ma011–1| 0.993      | 0.994       | 0.978       | 0.993       | ID          | 0.996       | 0.996       | 0.993       | 0.990       |
| Ma020–1| 0.996      | 0.998       | 0.981       | 0.994       | 0.996       | ID          | 1000        | 0.996       | 0.993       |
| Ma020–3| 0.996      | 0.998       | 0.981       | 0.994       | 0.996       | 1000        | ID          | 0.996       | 0.993       |
| AB016355| 0.993 | 0.995       | 0.985       | 0.991       | 0.993       | 0.996       | 0.996       | ID          | 0.996       |
| AB016347| 0.990 | 0.991       | 0.981       | 0.988       | 0.990       | 0.993       | 0.993       | 0.996       | ID          |
The symptoms of the disease described in the literature were identical with those presented in our study. The symptomatic leaves were covered with irregular spots with yellow haloes, which later changed into necrotic dark brown spots. Moreover, the incidence was much higher during the rainy season (Baharuddin 1997), comparably with observations in Poland.

Fig. 2 Results of maximum likelihood analysis of the 16S rRNA gene partial nucleotide sequences of the obtained isolates, in relation to both plant-pathogenic and non-pathogenic Pseudomonas species. The Pectobacterium carotovorum sequence (NR_025316.1) was used as outgroup to root the tree. The Polish P. syringae isolates are marked with black squares. BioEdit (v7.0.5.3) and Mega X software was used for multiple sequence alignments and the phylogenetic analysis, based on ClustalW and maximum likelihood algorithms implemented with BioEdit and Mega programs, respectively. Bootstrap values of 1000 pseudoreplicates were used for phylogenetic comparisons.

Similar to other plant hosts, the infection with P. syringae pv. mori has severe consequences for mulberry crops. The quality of mulberry leaves is of special importance because the leaves are used as a fodder for the silkworm B. mori, which has been domesticated for centuries and therefore is very sensitive to changes in breeding conditions and fodder. It is obvious that infection of mulberry leaves influences the growth and
development of silkworm larvae and ultimately the economical cocoon parameters (Y. Chen et al. 2018). That is why the proper breeding conditions of silkworms and the dietary components of mulberry leaves are so important for silk production, immune responses and resistance to pathogen infection of this valuable insect.

The systematic position of \textit{P. syringae} pv. mori is continuously changing. Multilocus sequence analyses confirm there are multiple species and pathovars of \textit{P. syringae} (Dillon et al. 2019; Yamamoto et al. 2000). According to recent studies involving DNA relatedness and ribotyping the following pathovars of \textit{P. syringae} are included in \textit{Pseudomonas amygdali}, namely \textit{tabaci, lachrymans, phaseolicola, mori}, among others (Haeyoung et al. 2015; Yamamoto et al. 2000); however, both species names are still accepted.

Despite extensive studies on management of \textit{P. syringae}, effective means of controlling this pathogen have not been established. Several methods of protection against \textit{P. syringae} have been tested on mulberry. For example, soil treatment with formaldehyde was tested against bacterial canker of white mulberry seedlings. Despite some efficiency, it turned out that such treated mulberry seedbeds should not be located near older mulberry trees because of its phytotoxic influence (Barbour et al. 2008). Also, attempts of developing an antimicrobial antioxidant from the plant source to treat plant pathogenic bacteria such as \textit{P. syringae} have been made. Aqueous

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{Fig_3.png}
\caption{Results of maximum likelihood analysis of the \textit{gyrB} gene partial nucleotide sequences of the obtained isolates, in relation to both plant-pathogenic and non-pathogenic \textit{Pseudomonas} species. The \textit{Pectobacterium carotovorum} sequence (FJ652727.1) was used as outgroup to root the tree. The Polish \textit{P. syringae} isolates are marked with black circles. BioEdit (v7.0.5.3) and Mega X software was used for multiple sequence alignments and phylogenetic analysis, based on ClustalW and neighbour joining algorithms implemented with BioEdit and Mega programs, respectively. Bootstrap values of 1000 pseudoreplicates were used for phylogenetic comparisons.}
\end{figure}
Fig. 4 Symptoms observed after spraying bacterial suspensions of *Pseudomonas syringae* on mulberry leaves. A-D – the pictures were aligned according to increasing symptoms development. Negative control: leaves treated with sterile distilled water (K-); Positive control: leaves treated with the *Pseudomonas syringae* pv. *syringae* 2238 strain (K+).
extracts of leaves of three Morus species (M. indica, M. latifolia, and M. laevigata) were tested to evaluate their antibacterial and antioxidant activity. Of the tested plants, the extracts of M. indica, and M. latifolia showed the highest antibacterial and antioxidant activity, and proved that the studied plant species are indeed important sources of pharmaceutical value (Joty et al. 2019). Nevertheless a specific plant protection compound based on those extracts is still not available commercially.

In summary, the studies we performed have proven P. syringae pv. mori to be the causal agent of mulberry bacterial blight in Poland. To our knowledge this is the first isolation, description and characterization of P. syringae affecting mulberry trees in Europe. This finding serves as a warning for phytosanitary services in Europe and for the sericulture industry that a new disease threat has appeared for the European mulberry crop.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

Research involving human participants and/or animals Not applicable. The research involved no human participants or animals.

Informed consent The research involved no human participants, and no animals so that the statement on the welfare of animals is not required.

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