Bacterial Metabolites Produced Under Iron Limitation Kill Pinewood Nematode and Attract Caenorhabditis elegans

Diogo Neves Proença*†, Thomas Heine2, Christoph H. R. Senges3, Julia E. Bandow3, Paula V. Morais1 and Dirk Tischler2,4

1 Department of Life Sciences and Laboratory of Environmental Microbiology of CEMMPRE, University of Coimbra, Coimbra, Portugal, 2 Environmental Microbiology, TU Bergakademie Freiberg, Freiberg, Germany, 3 Applied Microbiology, Faculty of Biology and Biotechnology, Ruhr University Bochum, Bochum, Germany, 4 Microbial Biotechnology, Faculty of Biology and Biotechnology, Ruhr University Bochum, Bochum, Germany

Pine Wilt Disease (PWD) is caused by Bursaphelenchus xylophilus, the pinewood nematode, and affects several species of pine trees worldwide. The ecosystem of the Pinus pinaster trees was investigated as a source of bacteria producing metabolites affecting this ecosystem: P. pinaster trees as target-plant, nematode as disease effector and its insect-vector as shuttle. For example, metals and metal-carrying compounds contribute to the complex tree-ecosystems. This work aimed to detect novel secondary metabolites like metallophores and related molecules produced under iron limitation by PWD-associated bacteria and to test their activity on nematodes. After screening 357 bacterial strains from Portugal and United States, two promising metallophore-producing strains Erwinia sp. A41C3 and Rouxiella sp. Arv20#4.1 were chosen and investigated in more detail. The genomes of these strains were sequenced, analyzed, and used to detect genetic potential for secondary metabolite production. A combinatorial approach of liquid chromatography-coupled tandem mass spectrometry (LC-MS) linked to molecular networking was used to describe these compounds. Two major metabolites were detected by HPLC analyses and described. One HPLC fraction of strain Arv20#4.1 showed to be a hydroxamate-type siderophore with higher affinity for chelation of Cu. The HPLC fraction of strain A41C3 with highest metal affinity showed to be a catecholate-type siderophore with higher affinity for chelation of Fe. LC-MS allowed the identification of several desferrioxamines from strain Arv20#4.1, in special desferrioxamine E, but no hit was obtained in case of strain A41C3 which might indicate that it is something new. Bacteria and their culture supernatants showed ability to attract C. elegans. HPLC fractions of those supernatant-extracts...
INTRODUCTION

The pine wilt disease (PWD) is one of the most devastating diseases of forests pine trees in the world, with tremendous ecological, environmental, and economic damage. Native to North America (United States and Canada), it has spread to Asia and Europe (Proença et al., 2017b). The only known causal agent is the nematode *Bursaphelenchus xylophilus*, the pinewood nematode (PWN; Nickle et al., 1981). PWN is transferred when an insect vector feeds on young shoots of pine trees and, if the tree is susceptible, develops the disease and starts the PWN-insect life cycle (Proença et al., 2017b). The ecosystem in which the PWD is developed is rather complex and comprises next to above mentioned tree-nematode-insect also adjacent microorganisms. Respectively, the endophytic microbiome of pine trees was recently characterized (Proença et al., 2017a; Alves et al., 2018), and the bacteria carried by PWN (Proença et al., 2010, 2014; Vicente et al., 2012) and insect vector were studied (Alves et al., 2016, 2018). Especially, the microbiome of the nematode seems directly linked to the tree-wilting (Alves et al., 2018). Further, the tree and nematode microbiome differ significant with respect to genus/species level. In addition, some of the relevant and more tree-related bacteria are endophytic opportunistic, living in soil and may enter the tree through the roots (Proença et al., 2017a). They are believed to be plant growth promoting rhizobacteria (PGPR) or do at least affect and thus interact with plants. They might not only promote the growth of the trees but also can control plant pests (Niranjan Raj et al., 2005). PGPR produce various compounds as for example small organic acids, phytohormones, antibiotics and siderophores among others. Metabolites like organic acids and siderophores interfere with metal or metalloid abundance and mobility in rhizosphere and surrounding environment. Especially the availability of metals, like iron (Fe$^{3+}$ or Fe$^{2+}$), was determined to be crucial within plant-microbe-nematode interactions (Ramamoorthy et al., 2001). It was reported that under iron limitation not lipopolysaccharides but siderophores as pseudobactin induce systemic resistance towards plant pests (Saikia et al., 2005; Romero et al., 2019). This response seems to be complementary to other defense mechanisms in dependence on the environmental conditions.

Metals are essential for life, acting as cofactors of enzymes and participating in central biological processes, as for example oxygen binding, activation and transport, photosynthesis, and within electron transport chains (Fones and Preston, 2013). The oxidation state of each metal is dependent on the environmental conditions that in turn results in its bioavailability for living organisms. Iron and copper are important metals for biological processes and its availability in soil influences plant growth as well as microorganisms (beneficial and pathogenic) (Zloch et al., 2016). Thus the redox environment and availability of certain metals and metalloids defines the ecosystem, especially in the rhizosphere and its vicinity (Zobel et al., 2005; McNear, 2013). In order to acquire certain metals and metalloids, organisms like microbes (e.g., the PGPR) and plants secrete metallophores which support the mobilization of poorly soluble metals (in case of Fe$^{3+}$ designated as siderophores) (Hider and Kong, 2010).

Metallophores have distinct biochemical characteristics and are divided into various families, i.e., hydroxamates, catecholates, carboxylates, and mixed types (Hider and Kong, 2010; Saha et al., 2013). Different types of siderophores have different metal binding capabilities and are often not restricted to iron but can scavenge different metals like copper, zinc, vanadium, gallium, and cadmium from the environment (Mehnert et al., 2017; Wiche et al., 2017; Schwabe et al., 2018). There are several biotechnological applications of siderophores, for example, enhancing of plant growth, acting as biological control agents (BCA), to perform bioremediation of environments contaminated with metals, to recover rare earth elements, and to act as antibiotic carriers or inhibitors of metalloenzymes (Ahmed and Holmström, 2014; Kurth et al., 2016). Plants are not able to assimilate iron in its predominant form in soil, ferric ion or Fe$^{3+}$, and produce phytosiderophores that mobilize Fe$^{3+}$ and are being taken up as Fe$^{3+}$-complexes (Ahmed and Holmström, 2014). Bacterial siderophores have been shown to be important in this interplay, increasing uptake of iron into tissues of the model plant *Arabidopsis thaliana* and improved plant growth (Vansuyt et al., 2007). On the other hand, bacterial siderophores are suggested to be pathogenicity factors because they compete for iron, limiting iron uptake by the plant (Gamalero and Glick, 2011; Wilson et al., 2016).

Overall it can be stated that metal limitation is stress for all the organisms of such an ecosystem described herein and the response siderophore production, and secretion can have various outcomes for the individuals. And as described above this can also affect the PWD-causing nematode and needs, therefore, to be investigated. The effects of siderophores towards the model nematode *Caenorhabditis elegans* is controversial. It was shown that bacterial siderophores from pseudomonads chelate iron and lead to death of *C. elegans* (Kirienko et al., 2013; Kang et al., 2018). However, siderophores like enterobactin caused an increase in iron levels in mitochondria, promoting the growth of *C. elegans* (Qi and Han, 2018).

Considering the large interest in the metabolic arsenal of environmental bacteria with biotechnological potential, the present work focused on the siderophores produced by wood of *Erwinia* strain A41C3, enriched with secondary metabolites such as siderophores, were able to kill pinewood nematode. These results suggest that metabolites secreted under iron limitation have potential to biocontrol *B. xylophilus* and for management of Pine Wilt Disease.

**Keywords:** metallophores, siderophore, nematode, *Erwinia*, *Rouxiella*, *Caenorhabditis elegans*, *Bursaphelenchus xylophilus*, secondary metabolites
colonizing and/or endophytic bacteria from the rhizosphere, where a complex community comprising plants and nematodes among others, interacts. Therefore, the present work aims to identify the siderophores produced by endophytic bacteria isolated associated with *Pinus* trees, in order to understand the role of bacterial siderophores using the model nematode *C. elegans* and verify their nematicidal activity towards the PWN. Two strains producing distinct siderophore patterns were chosen as model systems to investigate siderophore production, metal binding properties, and the biochemical properties of the siderophores, as well as their pathogenicity against nematodes.

**MATERIALS AND METHODS**

**Bacterial Strains, Culture Conditions and Biochemical Characterization**

Two hundred and seventy-one bacterial strains isolated as wood colonizers/endophytes of *P. pinaster* trees from Portugal (Proença et al., 2017a), 48 bacterial strains carried by PWN from Portugal (Proença et al., 2010), and 38 bacterial strains carried by PWN from United States (Proença et al., 2014) were evaluated for their production of siderophores on Iron-CAS agar plates (Schwyn and Neilands, 1987). The screening for bacterial endophytes and bacteria carried by PWN nematode from Portugal were re-evaluated in this study alongside with bacterial strains carried by PWN from United States for a confident result. The two best bacterial siderophores-producers candidates, with highest orange halo (positive for siderophore secretion) and lack of information of siderophores-producing species, strains Arv20#4.1 and A41C3, were selected for further production and characterization.

These two strains were grown in tubes with 5 ml of M9 media, supplemented with glucose (4% w/v), succinic acid (4 g/l), and Mg/Ca/B1/Goodie Mix (Esuola et al., 2016) for seven days. To investigate siderophore production and bacterial growth, the supernatant of each strain was obtained at days 0, 3, 5, and 7 and the OD<sub>600</sub> was determined at the same time points.

The phenotypes of strains Arv20#4.1 and A41C3 were analyzed focusing on the solubilization of phosphate and zinc, production of siderophores, and hydrolyzing activities including protease (casein), lipase (Tween 20, 40, 60 and 80), cellulase, and chitinase activities (Naveed et al., 2014). The ability to oxidize different carbon sources was assessed using Biolog GN2 MicroPlates incubated at 30°C. The results were recorded daily for up to 72 h using a MicroPlate reader (Tecan Infinite M200).

**Phylogenetic Analysis**

The strains Arv20#4.1 and A41C3 were grown on R2A agar media at 30°C for 48 h and its genomic DNA was extracted using the E.Z.N.A. Bacterial DNA Kit (Omega Bio-Tek, Norcross, GA, United States) according to the manufacturer’s instructions.

These strains were phylogenetically identified by 16S rRNA gene sequencing using primers 27F and 1525R as previously described (Rainey et al., 1992). All obtained sequences from bacterial isolates were compared with sequences available in the EMBL/GenBank database using BLASTN network services (Altschul et al., 1997) and with sequences available on the Eztaxon-e server<sup>1</sup> (Kim et al., 2012). Sequences were aligned within the SINA alignment service (Pruesse et al., 2012). Sequences were also checked for chimeric artifacts by using Mallard software (Ashelford et al., 2006). Sequences were included in 16S rRNA-based Living Tree Project (LTP) release 115 database<sup>2</sup> by parsimony implemented in the ARB software package version 5.5 (Ludwig et al., 2004). Phylogenetic dendrograms of sequences from this study and closest reference sequences were constructed by the Randomized Axelerated Maximum Likelihood (RAxML) method with GTRGAMMA model (Stamatakis, 2006) included in the ARB software (Ludwig et al., 2004).

**Genome Sequencing, Annotation and Siderophore Mining**

The draft genome sequencing of strains Arv20#4.1 and A41C3 was generated on an Illumina platform (MiSeq) as described previously (Retamal-Morales et al., 2018). To this end standard Illumina protocols and kits were employed. The extracted genomic DNA (see above) was used to prepare a genomic library using the NexteraXT DNA library kit. A quality check was performed using an Agilent Bioanalyzer 2100. The paired-end libraries obtained were sequenced employing v3-v4 chemistry on the MiSeq platform. The reads obtained were analyzed by applying FastQC, TrimGalore including the Phred20 quality filter, and SPAdes 3.5 for read assembly by using the GALAXY webtool<sup>3</sup>. The quality of the assembled genomes was accessed by means of Quast (Gurevich et al., 2013). The assembled genomes were annotated by using automatic pipelines of NCBI Prokaryotic Genomes Automatic Annotation Pipeline (PGAP)<sup>4</sup>, RAST (Aziz et al., 2008) and JGI servers (Huntemann et al., 2015; Chen et al., 2016, 2017).

Both annotated draft genomes were subjected to secondary metabolite gene cluster analysis by using the platform antiSMASH 3.0 (Weber et al., 2015), with specific interest for siderophores. Therefore, a BLAST analysis was performed for all protein sequences codified by each gene from siderophore gene cluster by using NCBI (Altschul et al., 1997) and UniProtKB/SwissProt and Uni-ProtKB/TrEMBL (Schneider et al., 2004) databases.

**Determination of Siderophore Production**

Supernatants obtained from cultures of strains Arv20#4.1 (bacterium carried by PWN from United States) and A41C3 (endophytic bacterium of *P. pinaster* from Portugal) were assessed for the siderophore activities by a modified CAS assay and a modified Csáky test (Esuola et al., 2016; Mehrert et al., 2017). EDTA and desferrioxamine B (DFOB) were used as positive controls. The modified CAS assay was used to determine

---

<sup>1</sup>http://eztaxon-e.ezbiocloud.net/

<sup>2</sup>http://www.arb-silva.de/projects/living-tree/

<sup>3</sup>https://usegalaxy.eu

<sup>4</sup>http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html
Fe, Ga, Cu, V, and Al chelating compounds in comparison to the same control reagents as for the Csáky test (Mehnert et al., 2017). Fe-, Ga-, Al-CAS solutions were prepared as follows (100 ml): 1.5 ml of 1 mM FeCl\textsubscript{3} × 6H\textsubscript{2}O (in 10 mM HCl) or 1.5 ml of 1 mM Ga\textsubscript{2}(SO\textsubscript{4})\textsubscript{3} or 1.5 ml of 1 mM AlCl\textsubscript{3} × 6 H\textsubscript{2}O was added to 7.5 ml of 2 mM aqueous CAS solution. Six ml of a 10 mM hexadecyltrimethylammonium bromide (CTAB) solution was added slowly. Subsequently MES buffer was added (10.67 g MES in 80 ml H\textsubscript{2}O with a pH of 5.6 by adjusting with 50% KOH). The mixture obtained was filled to a final volume of 100 ml with H\textsubscript{2}O. Cu-CAS solution was prepared as follows (100 ml): 2 ml of 10 mM CuCl\textsubscript{2}, 10.5 ml of 2 mM aqueous CAS solution, 2.5 ml of ddH\textsubscript{2}O and 85 ml of MES buffer at pH 5.7. V-CAS solution was prepared as follows (100 ml): 1.5 ml of 5 mM VCl\textsubscript{2}, 7.5 ml of 2 mM aqueous CAS solution, 6 ml of 10 mM CTAB and 85 ml of MES buffer at pH 4.1. Supernatants were mixed with Fe-CAS, Al-CAS, or V-CAS solution in a ratio 1:1 or were mixed with Ga-CAS or Cu-CAS solutions in a ratio 1:5. 33 mM of EDTA (in 10 mM HCl) and hydroxamate and catecholate siderophores were quantified as mentioned in previous section.

**LC-MS/MS Analysis**

Fractions from HPLC profiling with a significant CAS activity were analyzed by LC-MS/MS and fragment spectra were investigated by molecular networking. Samples were dried under reduced pressure and reconstituted in 1 ml of MS-grade methanol. To achieve a better solubility these samples were sonicated for 5 min. Mass spectrometric analysis was performed similarly as described earlier (Senges et al., 2018). Liquid chromatography (LC) was performed on a nanoACQUITY-UPLC system (Waters, Milford, MA, United States) with a Mixer Assy (Waters, zirc bead, inner cross section 1 mm, length 50 mm) and an AcquityUPLC HSS T3 column (Waters, core size 100 Å, particle size 1.8 μm, inner cross section dimension 1 mm, length 100 mm). 4.7 μl of each sample was injected and separated by means of a H\textsubscript{2}O/acetonitril (ACN) gradient with 0.1% formic acid (FA) and a flow rate of 25 μl min\textsuperscript{-1} (Table 1).

Subsequently, the mass spectrometry (MS) was performed with a Synapt G2-S HDMS\textsuperscript{E} system (Waters) with an ESI source operated in positive and negative mode and a TOF detector. Masses in the range of 50 to 3000 m/z were detected with 0.5 s per scan and leucine enkephalin injected as reference mass every 30 s. Mass spectrometry settings: lockspray capillary current 2.5 kV, capillary current 2.5 kV, cone voltage 30 V, source temperature 120°C, cone gas flow 60 l/h, flushing gas flow 550 l/h, with a temperature of 150°C. Fragmentation in MS/MS mode was conducted by collision induced diffraction with argon and a collision energy ramp of 10–25 V. Fragmentation was started when the intensity of a mass signal exceeded 6000 counts/s, finished after 6 s at max, and stopped prematurely if the intensity

| Time [min] | % ACN with 0.1% FA |
|------------|-------------------|
| 0.0        | 5.0               |
| 2.5        | 5.0               |
| 21.0       | 99.5              |
| 23.0       | 99.5              |
| 28.5       | 5.0               |
| 30.0       | 5.0               |

**Table 1** | Gradient used for LC-MS/MS analysis.

Frontiers in Microbiology | www.frontiersin.org 4 September 2019 | Volume 10 | Article 2166
Nematodes and eggs were centrifuged 1,300 g x 30 s at 22°C by vortexing every 2 min with final duration of 10 min. The addition of 2N NaOH and 1 ml of 3% sodium hypochlorite, followed by centrifugation in a total of 2 ml. It was added a fresh solution containing 0.5 ml of each bacterial supernatant, without removing any growth product. To test the nematicidal activity, 500 µl of each bacterial supernatant, with less than 0.06 O.D.600 nm, were incubated with 30 disinfected nematodes, for 48 h at 25°C. The assays were performed in triplicate. Nematodes were disinfected by sequential washes in sodium hypochlorite 0.1% (one wash, 1 min at 4°C) and in 1 ml sterilized water (two washes, 3 min at 4°C), followed by centrifugation. Last water (100 µl) was inoculated on R2A for control of the disinfection efficiency. The number of dead nematodes was assessed under the stereoscopic microscope. Nematodes were considered dead when linearized and not able to recover after being transferred to water. The controls, nematodes in CAA, were incubated under the same conditions.

Statistical Analysis
Chi-squared test was used to evaluate the significance of the differences in the chemotaxis of C. elegans with bacterial and supernatant samples. All differences were considered to be statistically significant for p < 0.05.

The significance of the differences, observed between the different nematicidal treatments towards B. xylophilus PtAS18 by using bacterial supernatants or HPLC fractions, was evaluated with two-way analysis of variance (ANOVA), using Bonferroni post-tests. The statistical analysis was performed with GraphPadPrism v5.0 for windows, GraphPad software, San Diego, CA, United States. All differences were considered to be statistically significant for p < 0.05.

Accession Numbers
The Whole Genome Shotgun project of strain Arv20#4.1 and A41C3 have been deposited at ENA under the accession numbers RQVU00000000 and RQVV00000000, respectively. The 16S rRNA gene sequences of the isolates, have been previously deposited in the NCBI GenBank database under the accession numbers according KF214948 (Proença et al., 2014) and KJ654833 (Proença et al., 2017a).

Accession links for metabolomic data sets are provided at: positive mode (http://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=bbd4f668b6224bd381617001be917381); negative mode (http://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=e3009a90a0954791ae9b747e727c6515).

C. elegans-Bacterial Strains/Supernatants Attraction Assay
Chemotaxis assays were performed according to Akduman et al. (2018) with some modifications. Briefly, 10 µl of bacterial suspensions obtained after 24 h of growth in M9 broth media supplemented with iron, at 30°C at 130 rpm, were placed 0.5 cm away from the edge of a 6 cm NGM agar plate. Ten µl of E. coli OP50 were added on the opposing side. Idealizing an equilateral triangle, 10 µl containing approximately 30 J3 stage nematodes were placed on the remaining vertex, with all spots at the same distance.

In the same line, 10 µl of bacterial supernatants, obtained from bacterial growth as mentioned above, was used to perform the chemotaxis attraction. Moreover, the bacterial strains were also grown in M9 broth media without supplementation of iron, and also its suspensions and its supernatants were tested in chemotaxis attraction of C. elegans N2.

Different combinations were performed to understand the attraction of the nematodes by different bacteria, its supernatants and if the influence of the presence of iron in M9 broth media also affects their attraction. M9 broth media with or without iron supplementation was also tested in chemotaxis assays. After 3h the number of nematodes were recorded in each test spot. All the assays were performed in triplicate.

Nematodes Survival Assays
The bacterial strains and their supernatants mentioned above were used to evaluate their effect on C. elegans N2 viability. The nematodes were observed and scored at 24 h, 48 h, and 72 h for with a dissecting microscope. On the other hand, the nematicidal activity of bacterial strains, its supernatants and HPLC fractions were evaluated towards the B. xylophilus PtAS18 (mixed life stages: J4 and adults) and towards C. elegans N2 (mixed life stages: L4 and adults) according to Paiva et al. (2013) with little modifications. Briefly, the bacterial suspensions obtained were centrifuged (20 min, 4°C, 13,000 rpm), in order to drastically reduce the number of cells, without removing any growth product. To test the nematicidal activity, 500 µl of each bacterial supernatant, with less than 0.06 O.D.600 nm, were incubated with 30 disinfected nematodes, for 48 h at 25°C. The assays were performed in triplicate. Nematodes were disinfected by sequential washes in sodium hypochlorite 0.1% (one wash, 1 min at 4°C) and in 1 ml sterilized water (two washes, 3 min at 4°C), followed by centrifugation. Last water (100 µl) was inoculated on R2A for control of the disinfection efficiency. The number of dead nematodes was assessed under the stereoscopic microscope. Nematodes were considered dead when linearized and not able to recover after being transferred to water. The controls, nematodes in CAA, were incubated under the same conditions.

Nematode Strains and Growth Conditions
The nematode wild type Caenorhabditis elegans N2, obtained from Caenorhabditis Genetic Center, Minneapolis, MN (United States) were propagated at 19.5°C on nematode growth agar medium (NGM) with Escherichia coli OP50 lawn (Brenner, 1974). After 3 days, nematodes were harvested by rinsing the plates with sterile M9 broth medium, transferred to 15 ml tubes and placed on the remaining vertex, with all spots at the same distance.

In the same line, 10 µl of bacterial supernatants, obtained after 24 h of growth in M9 broth media supplemented with iron, at 30°C at 130 rpm, were placed 0.5 cm away from the edge of a 6 cm NGM agar plate. Ten µl of E. coli OP50 were added on the opposing side. Idealizing an equilateral triangle, 10 µl containing approximately 30 J3 stage nematodes were placed on the remaining vertex, with all spots at the same distance.

In the same line, 10 µl of bacterial supernatants, obtained from bacterial growth as mentioned above, was used to perform the chemotaxis attraction. Moreover, the bacterial strains were also grown in M9 broth media without supplementation of iron, and also its suspensions and its supernatants were tested in chemotaxis attraction of C. elegans N2.

Different combinations were performed to understand the attraction of the nematodes by different bacteria, its supernatants and if the influence of the presence of iron in M9 broth media also affects their attraction. M9 broth media with or without iron supplementation was also tested in chemotaxis assays. After 3h the number of nematodes were recorded in each test spot. All the assays were performed in triplicate.

The nematode wild type Caenorhabditis elegans N2, obtained from Caenorhabditis Genetic Center, Minneapolis, MN (United States) were propagated at 19.5°C on nematode growth agar medium (NGM) with Escherichia coli OP50 lawn (Brenner, 1974). After 3 days, nematodes were harvested by rinsing the plates with sterile M9 broth medium, transferred to 15 ml tubes and placed on the remaining vertex, with all spots at the same distance.

In the same line, 10 µl of bacterial supernatants, obtained after 24 h of growth in M9 broth media supplemented with iron, at 30°C at 130 rpm, were placed 0.5 cm away from the edge of a 6 cm NGM agar plate. Ten µl of E. coli OP50 were added on the opposing side. Idealizing an equilateral triangle, 10 µl containing approximately 30 J3 stage nematodes were placed on the remaining vertex, with all spots at the same distance.

In the same line, 10 µl of bacterial suspensions obtained after 24 h of growth in M9 broth media supplemented with iron, at 30°C at 130 rpm, were placed 0.5 cm away from the edge of a 6 cm NGM agar plate. Ten µl of E. coli OP50 were added on the opposing side. Idealizing an equilateral triangle, 10 µl containing approximately 30 J3 stage nematodes were placed on the remaining vertex, with all spots at the same distance.

In the same line, 10 µl of bacterial suspensions obtained after 24 h of growth in M9 broth media supplemented with iron, at 30°C at 130 rpm, were placed 0.5 cm away from the edge of a 6 cm NGM agar plate. Ten µl of E. coli OP50 were added on the opposing side. Idealizing an equilateral triangle, 10 µl containing approximately 30 J3 stage nematodes were placed on the remaining vertex, with all spots at the same distance.

In the same line, 10 µl of bacterial suspensions obtained after 24 h of growth in M9 broth media supplemented with iron, at 30°C at 130 rpm, were placed 0.5 cm away from the edge of a 6 cm NGM agar plate. Ten µl of E. coli OP50 were added on the opposing side. Idealizing an equilateral triangle, 10 µl containing approximately 30 J3 stage nematodes were placed on the remaining vertex, with all spots at the same distance.

In the same line, 10 µl of bacterial suspensions obtained after 24 h of growth in M9 broth media supplemented with iron, at 30°C at 130 rpm, were placed 0.5 cm away from the edge of a 6 cm NGM agar plate. Ten µl of E. coli OP50 were added on the opposing side. Idealizing an equilateral triangle, 10 µl containing approximately 30 J3 stage nematodes were placed on the remaining vertex, with all spots at the same distance.

In the same line, 10 µl of bacterial suspensions obtained after 24 h of growth in M9 broth media supplemented with iron, at 30°C at 130 rpm, were placed 0.5 cm away from the edge of a 6 cm NGM agar plate. Ten µl of E. coli OP50 were added on the opposing side. Idealizing an equilateral triangle, 10 µl containing approximately 30 J3 stage nematodes were placed on the remaining vertex, with all spots at the same distance.

In the same line, 10 µl of bacterial suspensions obtained after 24 h of growth in M9 broth media supplemented with iron, at 30°C at 130 rpm, were placed 0.5 cm away from the edge of a 6 cm NGM agar plate. Ten µl of E. coli OP50 were added on the opposing side. Idealizing an equilateral triangle, 10 µl containing approximately 30 J3 stage nematodes were placed on the remaining vertex, with all spots at the same distance.
RESULTS

Siderophore Screening, Bacterial Phylogeny and Biochemical Characterization

In this study, 357 bacterial strains were tested and belonging to 52 genera according to 16S rRNA gene analysis (Proença et al., 2010, 2014, 2017a). Numerous bacterial strains were positive on CAS agar medium that serves to detect siderophores: 103 endophytic strains (38.0%), 39 strains carried by *B. xylophilus* from Portugal (81.2%) and 20 strains carried by *B. xylophilus* from United States (52.6%) (Supplementary Figure S1). Among those, two isolates showed a distinct siderophore production on CAS agar plates and have been chosen for further study: strains Arv20#4.1 and A41C3.

According to neighbor-joining and maximum-likelihood phylogenetic trees, the closest relatives to strain Arv20#4.1 were *Rouxiella chamberiensis* (99.6% sequence similarity), *R. silvae* (99.6%), and *R. badensis* (99.6%). *Erwinia endophytica* (98.1%), *Pantoea dispersa* (97.7%), and *E. psidii* (97.6%) were the closest relatives to strain A41C3. Thus, the best two bacterial isolates with higher biotechnological potential analyzed in this study belong to the family *Enterobacteriaceae*, and were *Erwinia* sp. A41C3 (endophyte from Portuguese pines) and *Rouxiella* sp. Arv20#4.1 (bacterium isolated as carried by PWN from United States) (Supplementary Figure S2).

Biochemical characteristics of strains Arv20#4.1 and A41C3 are summarized in Table 2. Both strains were able to solubilize phosphate and zinc, produce siderophores, degrade *Twee* 60 and carboxymethyl cellulose (CMC), and produced catalases; but were negative for protease production, chitin, *Twee* 40 and *Twee* 80 degradation. Both strains oxidized 43 carbon sources in GN2 MicroPlate (and more carbon sources were weakly oxidized). Only the strain A41C3 was able to oxidize m-inositol, β-hydroxybutyric acid, α-keto butyric acid, α-keto glutaric acid, sebacic acid, succinic acid, L-prolylglutamic acid, γ-amino butyric acid in GN2 MicroPlates (Table 2).

Genome Analysis Showed (Novel) Siderophore Gene Clusters

The draft genome sequence of strains Arv20#4.1 contained 29 contigs, totaling 5.37 Mbp in size with a mapped coverage of 547-fold of the genome. The G + C content was 53.0 mol%. The genome encoded 4,936 putative coding sequences (CDSs) (Table 3). The draft genome sequence of strain Arv20#4.1 contained loci for three ribosomal RNAs and 72 tRNAs (Table 3).

In case of strain A41C3, the draft genome sequence contained 61 contigs, totaling 4.22 Mbp in size with a mapped coverage of 792-fold of the genome. The G + C content was 51.6 mol%. The genome encoded 3,953 putative CDSs (Table 3). The draft genome sequence of strain A41C3 contained loci for seven ribosomal RNAs and 57 tRNAs (Table 3). A general overview of the biological subsystems obtained using RAST for both strains is summarized in Table 4.

By using the webtool antiSMASH, it was possible to identify the gene cluster for secondary metabolite, especially siderophore, production close to desferrioxamine, in particular the operon that is composed by decarboxylase, monooxygenase, and acetyltransferase in strain Arv20#4.1 (Table 5 and Supplementary Figure S3A). For strain A41C3, it was possible

| Table 2 | Differential characteristics between strains Rouxiella sp. Arv20#4.1 and Erwinia sp. A41C3. |
|---------|--------------------------------------------------------------------------------------------------|
| Characteristic | Arv20#4.1 | A41C3 |
| Phosphate solubilization | W | + |
| Siderophores | +++ | ++ |
| Tween 20 | ++ | – |
| CMO degradation | + | – |
| Catalase | + | ++ |
| **BIOLOG** | | |
| Tween 80 | W | + |
| M-Inositol | – | + |
| D-Sorbitol | + | W |
| Turanose | W | + |
| Pyruvic Acid Methyl Ester | – | W |
| Citric Acid | W | + |
| Formic acid | + | W |
| D-Glucosaminic acid | W | + |
| α-Hydroxybutyric acid | W | + |
| β-Hydroxybutyric acid | – | + |
| P-Hydroxy Phenylacetic acid | – | W |
| α-Keto butyric acid | – | + |
| α-Keto glutaric acid | – | + |
| α-Keto valeric acid | – | W |
| Malonic acid | W | + |
| Quinic acid | W | + |
| Sebacic acid | – | + |
| Bromosuccinic acid | W | + |
| Succinic acid | – | + |
| L-Alaninamide | – | W |
| D-Alanine | + | W |
| D-Glutamic acid | W | + |
| Glycol-L-aspartic acid | W | + |
| L-Proline | W | + |
| L-Pyrogulotic acid | W | + |
| L-Theonine | W | + |
| γ-Amino butyric acid | – | + |
| Urocanic acid | W | + |

Both strains were positive on solubilization of zinc, degradation of Tween 60, and negative on production of proteases, degradation of *Twee* 40, *Twee* 80 and chitin. Both strains oxidized *Twee* 40, N-acetyl-D-glucosamine, adonitol, L-arabinose, D-arabitol, D-cellulose, erythritol, D-fructose, L-fucose, D-galactose, gentiobiose, α-D-glucose, m-inositol, α-D-lactose, lactulose, D-maltose, D-mannitol, D-mannose, D-melibiose, b-methyl-D-glucoside, D-psicose, D-raffinose, L-rhamnose, D-sorbitol, sucrose, D-trehalose, turanose, xylitol, pyruvic acid methyl ester, succinic acid mono-methyl-ester, acetic acid, cis-aconitic acid, citric acid, formic acid, D-galactonic acid, D-galacturonic acid, D-gluconic acid, D-glucosaminic acid, D-glucuronic acid, α-hydroxybutyric acid, β-hydroxybutyric acid, γ-hydroxybutyric acid, α-phosphoethanoic acid, itaconic acid, α-keto butyric acid, α-keto glutaric acid, α-keto valeric acid, D,L-lactic acid, malonic acid, propionic acid, quinic acid, D-saccharic acid, sebacic acid, succinic acid, bromosuccinic acid, succinic acid, glucuronamide, L-alaninamide, D-alanine, L-alanine, L-alanyl-glycine, L-asparagine, L-aspartic acid, L-glutamic acid, glycyl-L-aspartic acid, glycyl-L-glutamic acid, L-histidine, L-hydroxyproline, L-leucine, L-ornithine, L-phenylalanine, L-proline, L-pyroglutamic acid, D-serine, L-serine, L-threonine, D,L-carnitine, γ-amino butyric acid, uric acid, inosine, uridine, thymidine, phenylalanine, putrescine, 2-aminoethanol, 2,3-butanediol, glycerol, D,L-α-glycerol phosphate, α-D-glucose-1-phosphate, D-glucose-6-phosphate, +, Positive; W, weakly positive; –, negative.
to identify multiple genes in the gene cluster for siderophore production (Table 6 and Supplementary Figure S3B). However, it was not possible to predict which siderophore is produced. There are several putative siderophore biosynthesis clusters identified by antiSMASH that could be attributed to strain A41C3 like as turnerbactin, griseobactin among others.

Production and Biochemical Characterization of Siderophores

Both bacterial strains reached maximum of optical density after 3 days of incubation in 10 ml tubes containing 5 ml of M9 modified medium. Siderophore production by both strains depended on days of cultivation, but both supernatants showed activity for hydroxamate and catecholate-type siderophores (Figure 1). Strain Arv20#4.1 produced mainly hydroxamate-type siderophores (25 µM of NH₂OH equivalents) while strain A41C3 produced mainly catecholate-type siderophores (70 µM of DHBA equivalents) and showed more affinity to copper or to iron, respectively.

After three days, the growth medium of strain Arv20#4.1 had a pH of 9 while, that of A41C3 a pH of 5. At this moment siderophore comprising supernatants were harvested and processed by ion exchange material to obtain an enriched siderophore pool for each strain. After concentration by evaporation, siderophores were enriched by HPLC and 30 fractions per strain were obtained (Figure 2). Both strains showed ability to produce hydroxamate and catecholate-type siderophores but with different concentrations. The best metal affinity was identified in an HPLC fraction (1.25) of supernatant of strain Arv20#4.1 for a hydroxamate-type siderophore (92 µM of NH₂OH equivalents) with higher affinity for Cu (Figure 2B). On the other hand, the best supernatant HPLC fraction (3.26) of strain A41C3 contained a catecholate-type siderophore (309 µM of DHBA equivalents) with higher affinity for Fe (Figure 2D).

Siderophore Identification by Mass Spectrometry

The collected compounds in HPLC fractionations were characterized by using LC-MS. Fragment spectra were analyzed by molecular networking, which allows sorting of spectra according to their similarity that is ultimately based on the structural similarity of molecules analyzed (Wang et al., 2016). Guided by the genome-based biosynthetic potential mass spectrometric data were compared with literature and databases to identify the enriched molecules (Senges et al., 2018).

As indicated by its desferrioxamine biosynthetic gene cluster, Arv20#4.1 produces a set of desferrioxamines (Figure 3A). These were detected in both, positive and negative mode from the LC-MS procedure (Supplementary Figure S4). As the eponymous compound for this molecular subnetwork the hydroxamate siderophore desferrioxamine E was identified
TABLE 5 | Secondary metabolite clusters identified in strain Rouxiella sp. Arv20#4.1 with antiSMASH3.0.

| Cluster | Type | Start | Stop | Similarity to known cluster | MIBiG BGC-ID |
|---------|------|-------|------|------------------------------|--------------|
| 1       | Hserlactone | 87033 | 107683 | – | – |
| 2       | Cf_fatty_acid | 446284 | 467504 | – | – |
| 3       | Arylpolyene | 767129 | 821369 | APE Ec biosynthetic gene cluster (68% of genes show similarity) | BGC0000836_c1 |
| 4       | Cf_putative | 25500 | 34695 | Emulsan biosynthetic gene cluster (9% of genes show similarity) | BGC0000760_c1 |
| 5       | Cf_fatty_acid-Cf_saccharide | 24604 | 60942 | Lipopolysaccharide biosynthetic gene cluster (18% of genes show similarity) | BGC0000776_c1 |
| 6       | Cf_putative | 27900 | 36897 | – | – |
| 7       | Cf_putative | 89005 | 95978 | – | – |
| 8       | Cf_fatty_acid | 165825 | 186739 | – | – |
| 9       | Cf_saccharide | 485089 | 506842 | S-layer glycan biosynthetic gene cluster (20% of genes show similarity) | BGC0000794_c1 |
| 10      | Cf_fatty_acid | 532454 | 553725 | – | – |
| 11      | Cf_putative | 586070 | 593059 | Polysaccharide B biosynthetic gene cluster (6% of genes show similarity) | BGC0001411_c1 |
| 12      | Cf_putative | 712377 | 716502 | – | – |
| 13      | Cf_putative | 103014 | 112356 | – | – |
| 14      | Cf_putative | 283336 | 295617 | – | – |
| 15      | Siderophore | 270474 | 282858 | Desferrioxamine B biosynthetic gene cluster (60% of genes show similarity) | BGC0000941_c1 |
| 16      | Cf_putative | 164758 | 170600 | – | – |
| 17      | Cf_saccharide | 97264 | 131452 | Capsular polysaccharide biosynthetic gene cluster (33% of genes show similarity) | BGC0000731_c1 |

NRPS, Non-ribosomal peptide synthetase cluster; Cf, ClusterFinder.

TABLE 6 | Secondary metabolite clusters identified in strain Erwinia sp. A41C3 with antiSMASH3.0.

| Cluster | Type | Start | Stop | Similarity to known cluster | MIBiG BGC-ID |
|---------|------|-------|------|------------------------------|--------------|
| 1       | Cf_putative | 284735 | 289620 | Polysaccharide B biosynthetic gene cluster (8% of genes show similarity) | BGC0001411_c1 |
| 2       | Cf_putative | 15697 | 24996 | – | – |
| 3       | Cf_putative | 68401 | 78206 | – | – |
| 4       | Thiopeptide | 210237 | 236598 | O-antigen biosynthetic gene cluster (14% of genes show similarity) | BGC0000781_c1 |
| 5       | Cf_fatty_acid | 347964 | 368917 | Svaricin biosynthetic gene cluster (6% of genes show similarity) | BGC0001382_c1 |
| 6       | Cf_putative | 72078 | 80802 | – | – |
| 7       | Cf_fatty_acid | 1 | 11819 | – | – |
| 8       | Cf_putative | 18987 | 28258 | Emulsan biosynthetic gene cluster (9% of genes show similarity) | BGC0000760_c1 |
| 9       | Cf_fatty_acid | 1 | 7355 | – | – |
| 10      | Cf_fatty_acid -Ladderane | 35473 | 78034 | – | – |
| 11      | Cf_fatty_acid | 41264 | 62484 | – | – |
| 12      | Cf_saccharide | 70023 | 92437 | – | – |
| 13      | Cf_saccharide | 57635 | 93206 | Lipopolysaccharide biosynthetic gene cluster (60% of genes show similarity) | BGC0000779_c1 |
| 14      | Hserlactone | 73942 | 94574 | – | – |
| 15      | Cf_saccharide | 30591 | 54347 | – | – |
| 16      | Cf_saccharide | 103519 | 141097 | Stewartan biosynthetic gene cluster (85% of genes show similarity) | BGC0000763_c1 |
| 17      | Cf_saccharide-NRPS | 131937 | 194079 | Turnerbactin biosynthetic gene cluster (30% of genes show similarity) | BGC0000451_c1 |
| 18      | NRPS | 271826 | 315746 | – | – |
| 19      | Cf_putative | 129079 | 138254 | – | – |
| 20      | Cf_putative | 93961 | 107030 | – | – |
| 21      | Cf_putative | 143921 | 152024 | – | – |
| 22      | Butyrolactone | 192718 | 198822 | – | – |

NRPS, Non-ribosomal peptide synthetase cluster; Cf, ClusterFinder.

by comparison with reference spectra described in Senges et al. (2018) and through manual mapping of the fragment spectrum onto the molecular structure (Figure 3B). As sortation of fragment spectra by molecular networking allows sortation into structurally related families, further compounds in the desferrioxamine subnetwork are likely desferrioxamine E analog. This is supported by five further desferrioxamine-like siderophores, identified by their complete masses (Supplementary Table S1).

Of the 421 and 1,034 nodes in the molecular network of A41C3, originating from positive and negative mode respectively, none could be identified. One fragment spectrum showed similarity to a database entry of a desferrioxamine, but its characteristics (parent and fragment masses) were unlike those...
of typical desferrioxamines. Iron chelating and biological activity are likely to originate from different, yet unknown compounds. Respectively, an overview on siderophore activity of fractions with corresponding signals in mass spectrometry are provided in Figure 2 and Supplementary Table S2.

C. elegans Attracted by Siderophore-Producing Bacteria

Both bacteria, after 24 h growth, showed higher OD$_{600}$ in the M9 growth media supplemented with iron than in low-iron medium. This indicates iron limitation creates stress and should induce a metabolic response as shown above for siderophore production. Strain Arv20#4.1 reached an OD$_{600}$ of 2.06 when growth medium was supplemented with iron and 0.84 when growth medium was deprived of iron. In the same way, strain A41C3 showed OD$_{600}$ of 2.19 when growth medium was supplemented with iron and 0.54 when growth medium was deprived of iron.

C. elegans N2 was attracted by strains Arv20#4.1 and A41C3 and by their supernatants (Figure 4). The M9 broth media supplemented or not supplemented with iron did not attract nematodes. The E. coli OP50 was used as control and nematodes showed preference for their regular food source in comparison with both strains but without statistical significance. The nematodes were significantly attracted by strain A41C3 (+ Fe) compared to Arv20#4.1 (+ Fe), but no significant attraction of nematodes was observed when both strains were grown in M9 broth medium without iron supplementation. These differences might be due to differences in the metabolites produced and maximum cell density reached (OD$_{600}$). The supernatant of Arv20#4.1 with iron in the growth medium showed significant attraction of nematodes when compared with same strain supernatant without iron. The same was observed when supernatant of Arv20#4.1 (+ Fe) compared with supernatant of A41C3 supplemented with iron (Figure 4). On the other hand, supernatant of Arv20#4.1 without iron showed significant attraction of nematodes when compared with supernatant of A41C3 without supplementation of iron (Figure 4).

Nematicidal Activity

The herein investigated bacteria and their supernatants did not show nematicidal activity against C. elegans N2 evaluated at 24 h, 48h, and 72h incubation.

In the same line, the bacterial supernatants and fractions of both strains containing siderophores were tested against B. xylophilus. It was possible to observe significant nematicidal activity for HPLC fractions of the samples A-G against B. xylophilus PtAS18 (Figure 5). The sample G, containing HPLC fractions 3.25 + 3.26 + 3.27 + 3.28, showed the strongest nematicidal activity that includes combination of HPLC fractions 25, 26, 27, 28 (equal ratio) of the strain A41C3. The compounds present in these HPLC fractions, with nematicidal activity, showed similarity to a number of database entries with respect to the MS analysis. However, an identification was impossible so far. Therefore, the amounts were too little to obtain a relevant fragmentation pattern or even NMR data after iron removing.
Proença et al. Bacterial Metabolites Kill the Pinewood Nematode

FIGURE 2 | Purification and characterization of the siderophores in supernatant of strains Arv20#4.1 (A,B) and A41C3 (C,D) (A,C) The siderophores were enriched by HPLC using a C18-RP column (1 ml; Thermo, Aquabt). Equilibrated in 28% methanol (buffer A) and 72% ddH2O with 0.1% TFA (buffer B). The elution from the column was performed as described and after 8 min, a 4-min linear gradient 28–95% buffer A followed by 3 min 95% buffer A was applied before going back to initial conditions (total time of elution 15 min at 1 ml/min flow rate). (B,D) Characterization of 30 fractions for Fe, Ga, Cu, V, and Al-CAS activities and quantitation of hydroxamate and catecholate-type siderophores. The best supernatant HPLC fraction of strain Arv20#4.1 containing hydroxamate-type siderophores (92 µM of \( \text{NH}_2\text{OH} \) equivalents) most actively chelated Cu (B). The best supernatant HPLC fraction of strain A41C3 contains catecholate-type siderophores (309 µM of DHBA equivalents) with higher affinity for Fe (D).

| Fraction (t-min) | Fe\(^+\) | Ga\(^+\) | V\(^+\) | Cu\(^+\) | Al\(^+\) | Hydroxamate\(^a\) | Catecholate\(^a\) |
|-----------------|--------|--------|--------|--------|--------|----------------|----------------|
| 3.1 (0.0)       | -      | -      | 32.5   | 3.6    | -      | -              | -              |
| 8.2 (0.9)       | -      | -      | 34.8   | 8.4    | -      | -              | -              |
| 11.2 (1.8)      | -      | -      | 34.3   | 3.1    | -      | -              | -              |
| 14.1 (2.7)      | -      | 0.1    | 34.8   | 8.5    | -      | -              | -              |
| 15.5 (3.6)      | -      | 0.2    | -      | 13.8   | -      | -              | -              |
| 3.6 (2.5)       | -      | 0.8    | 31.5   | 8.7    | -      | -              | -              |
| 8.6 (3.4)       | 1.4    | 0.7    | 44.4   | 12.7   | -      | -              | -              |
| 8.4 (3.4)       | -      | 1.4    | 40.7   | 9.2    | -      | -              | -              |
| 9.5 (4.0)       | -      | 0.4    | 27.7   | 4.5    | -      | -              | -              |
| 12.0 (4.5)      | -      | 0.4    | 37.5   | 2.5    | -      | -              | -              |
| 3.4 (5.0)       | -      | 0.6    | 37.9   | 6.1    | -      | -              | 123.7          |
| 3.1 (5.3)       | 1.8    | 0.5    | 39.3   | 6.2    | -      | -              | 123.7          |
| 3.1 (6.0)       | 1.9    | 0.4    | 40.2   | 6.3    | -      | -              | 7.8            |
| 3.4 (6.5)       | 6.5    | 2.5    | 46.8   | 5.9    | -      | -              | 12.7           |
| 3.4 (7.0)       | 35.2   | 15.9   | 45.7   | 9.7    | -      | -              | 123.7          |
| 8.4 (7.6)       | 49.2   | 4.7    | 48.0   | 8.2    | -      | -              | 123.7          |
| 3.2 (7.8)       | 35.0   | 2.1    | 47.7   | 2.3    | -      | -              | 24.9           |
| 8.8 (8.4)       | 7.9    | 1.0    | 47.0   | 2.5    | -      | -              | 12.7           |
| 3.1 (9.0)       | 0.7    | 0.8    | 45.0   | 4.4    | -      | -              | -              |
| 3.2 (9.5)       | -      | 0.6    | 38.9   | 6.0    | -      | -              | -              |
| 3.2 (10.0)      | -      | 0.5    | 35.5   | 6.7    | -      | -              | -              |
| 3.2 (10.5)      | -      | 0.4    | 44.4   | 4.9    | -      | -              | -              |
| 3.2 (11.0)      | -      | 0.5    | 36.5   | 6.6    | -      | -              | -              |
| 3.2 (11.5)      | 5.4    | 0.6    | 43.5   | 8.1    | -      | -              | -              |
| 3.2 (12.0)      | 40.1   | 7.4    | 30.1   | 10.7   | 5.7    | -              | 26.0           |
| 3.2 (12.5)      | 96.7   | 62.0   | 18.6   | 29.6   | -      | -              | 309.2          |
| 3.2 (13.0)      | 176.1  | 12.7   | 11.7   | 10.4   | 12.2   | 7.3           | 150.6          |
| 3.2 (13.5)      | 65.6   | 2.9    | -      | 3.8    | -      | -              | -              |
| 3.2 (14.0)      | -      | 0.5    | -      | -      | -      | -              | -              |
| 3.2 (14.5)      | -      | -      | -      | -      | -      | -              | -              |

* [DFB equivalents] (µM); \(^a\) [NH\(_2\)OH equivalents] (µM); \(^b\) [DHBA equivalents] (µM); -, negative.
DISCUSSION

Bacteria are important in the ecosystem of P. pinaster trees as they are involved in key activities and interactions such as production of proteases, lipases, cellulases, secretion of metabolites for signaling and auxins (Proença et al., 2010, 2017a,b; Vicente et al., 2012). They are further involved in solubilization of phosphate and zinc and mobilization of metals and metalloids by the production of siderophores (Proença et al., 2010, 2017a,b). Among those activities, (secondary) metabolite production and release by microbes represents an important trigger in the ecosystems. This is often a response towards a limitation of resources within such an ecosystem. Here, we focused especially on metallophores produced under metal (iron) limitation and known to have species overlapping activities (Hider and Kong, 2010; Wiche et al., 2017; Senges et al., 2018). Metallophores are known to be part of the arsenal of plant-growth promoting (rhizo)bacteria and to interact with other microorganisms, nematodes, higher animals, and plants within the ecosystem (Mercado-Blanco et al., 2018). Metabolites like siderophores play important roles in ecosystems to chelate metallic micronutrients essential for microorganisms and plants. However, siderophores have been recognized to have a key role in virulence of pathogenic organisms (Soutar and Stavrínides, 2018) and thus its screening is important.

Species belonging to the family Enterobacteriaceae are known to grow in soil next to plants, to engage in plant-microbe interactions, and thus are part of a complex network in the rhizosphere. Strains of the genus Rouxiella have been isolated from peat bog soil (Le Fléche-Matéos et al., 2017) and are recognized to produce biosurfactants (Kügler et al., 2015). Strains belonging to both genera Erwinia and Pantoea have been demonstrated to produce hydroxamate siderophore desferrioxamine E among others that supports the iron acquisition by the strains. This ability favors host colonization and it is recognized as a factor of pathogenicity (Dellagi et al., 2009).

Here, we found that the endophytic Erwinia strain A41C3 produces a catecholate-type siderophore based on analytical methods and bioinformatics genome annotation pipeline. The latter hints towards turnerbactin but this was not verified by LC-MS/MS and thus, these compounds remain enigmatic and await structural identification. Erwinia species are known to produce achromobactin or aerobactin (Soutar and Stavrínides, 2018), two hydroxamate-type siderophores, that have not yet been described to produce catecholate-type metallophores like turnerbactin-related compounds. This will be investigated in more detail in the near future.

In this study, we found that PWN-carried Rouxiella strain Arv20#4.1 produces desferrioxamine-like compounds based on analytical methods and also predicted from the genome (Figure 3 and Supplementary Figure S2). This was verified by LC-MS/MS confirming the identification of desferrioxamine-like structures, with desferrioxamine E (DFOE) as major metabolite. This is the first time that siderophore production was described in more detail by a strain of the genus Rouxiella and the respective siderophore was identified as DFOE. The HPLC-enriched fraction of DFOE and related molecules (Figure 2) showed affinity to various metals and metalloids as described previously for other desferrioxamines obtained from soil bacteria (Mehnert et al., 2017; Wiche et al., 2017; Schwabe et al., 2018).
As described above both strains, A41C3 and Arv20#4.1, produced different siderophores which have different properties. As various metals and metalloids may play a role in the rhizosphere ecosystem it is of interest to verify the capability of the siderophores to bind various types of ions (Mehnert et al., 2017; Wiche et al., 2017; Schwabe et al., 2018). Therefore, we decided to choose a selection of ions which can form a CAS-complex and thus allow to show and to discriminate affinity of siderophores for Fe, Ga, Cu, V, and Al (Figure 2). The siderophore of strain Arv20#4.1 is a hydroxamate-type siderophore and has a high affinity for Cu. The one of strain A41C3 is a catecholate-type siderophore and has a high affinity for Fe.

**Caenorhabditis elegans** N2 were attracted by both strains and by their supernatants. This attraction of this nematode through different bacterial species has been recognized in several studies with other bacterial strains (Choi et al., 2016). No nematicidal activity against **C. elegans** was demonstrated by either species, by bacterial supernatants, or by HPLC fractions containing siderophores. Monitoring food behavior of **C. elegans** is a simple approach to identify potential bacterial virulence (Lewenza et al., 2014). Considering that the biological role of siderophores is to ensure the acquisition of iron necessary for metabolism, their production is expected to be lower in bacteria growing in iron-supplemented media. This was the case, however, the nematode behavior was similar in both supernatants. **C. elegans** preferred...
bacterial supernatants to bacterial cells, independently of the bacterial growth conditions. This behavior is in congruence with the fact that bacterial supernatants and siderophores do not kill C. elegans and may be used by the nematode as food sources. The type of siderophores, and probably the concentration, might be crucial for nematicidal activity as it was demonstrated in pathogenicity of P. aeruginosa by its siderophore pyoverdin (Kirienko et al., 2013). On the other hand, the nematode behavior seems to indicate that Rouxia sp. Arv20#4.1 is more virulent than Erwinia sp. A41C3, using virulence factors different from siderophores. Both bacterial species are gamma-proteobacteria reported to be pathogenic (Paiva et al., 2013).

The effect of enriched siderophore fractions on B. xylophilus behavior was evaluated for the first time. The isolate Arv20#4.1 originating from PWN did produce mainly desferrioxamines which had no effect on B. xylophilus. Thus it might be reasoned that desferrioxamines interfere with the mobility of metals but do not affect other organisms of this ecosystem to a large extent. However, this nematode showed to be highly susceptible to supernatants containing siderophores and other metabolites produced by the endophytic Erwinia sp. A41C3 under iron limitation. Here, the highest nematicidal activity was determined from the fractions comprising highest siderophore titer, respectively. This seems to be a clear correlation and needs to be elucidated with pure compounds in more detail. Siderophores produced by entomopathogenic bacteria (Hirschmann et al., 2017), endophytic plant growth-promoting bacteria (Tashioshnoei et al., 2017), and rhizospheric bacteria (Khan et al., 2016) have been related with the pathogenic activity of these bacteria against insects, plant pathogenic bacteria and root-knot nematode, respectively. The role of this Erwinia strain when part of the endophytic bacterial community needs to be explored. In spite of that, their metabolites produced under iron limitation should be investigated further as they might prove useful in biocontrol B. xylophilus and management of Pine Wilt Disease.

**DATA AVAILABILITY STATEMENT**

The Whole Genome Shotgun project of strains Arv20#4.1 and A41C3 has been deposited at ENA under the accession numbers RQVU00000000 and RQVV00000000, respectively. The 16S rRNA gene sequences of the isolates have been previously deposited in the NCBI GenBank database under the accession numbers KF214948 (Proença et al., 2014) and KJ654833 (Proença et al., 2017a). Accession links for metabolomic data sets are provided at: positive mode (http://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=bbd4f68b6224bd381617001be917381); negative.
mode (http://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=e3009a90a0954791ae9b747e727c6515).

AUTHOR CONTRIBUTIONS

DP, PM, and DT designed the experiments and wrote the manuscript. DP did all the experiments. TH did the genome sequencing and analysis. CS performed the mass spectrometry sequencing and analysis. DP, PM, and DT designed the experiments and wrote the manuscript. All authors approved the manuscript.

FUNDING

This work was supported by the CEMMPRE and Fundação para a Ciência e a Tecnologia (FCT) under the projects UID/EMS/00285/2013, PTW-PTDC/AAG-REC/3839/2014, and ERA-MIN/0002/2015. This work was also supported by the COST Action FP1305 – Biolink: Linking belowground biodiversity and ecosystem function in European forests. DT and TH were supported by a Junior Research Grant from the German Federal Ministry of Education and Research (BakSolEx 033R147). JB acknowledges the funding from the German Research Foundation (BA 4193/6-1) and German federal state of North Rhine-Westphalia (NRW) for the mass spectrometer (Forschungsgroßgerät der Länder).

ACKNOWLEDGMENTS

We thank Dr. Luis Fonseca for providing the pinewood nematodes used in this study.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2019.02166/full#supplementary-material

REFERENCES

Ahmed, E., and Holmström, S. J. M. (2014). Siderophores in environmental research: roles and applications. Microb. Biotechnol. 7, 196–208. doi: 10.1111/1751-7915.12117
Akduman, N., Rödelperger, C., and Sommer, R. J. (2018). Culture-based analysis of Pristionchus-associated microbiota from beetles and figs for studying nematode-bacterial interactions. PLoS One 13:e0198018. doi: 10.1371/journal.pone.0198018
Altschul, S. F., Madden, T. L., Schäffer, A. A., Zhang, J., Zhang, Z., Miller, W., et al. (1997). Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res. 25, 3389–3402. doi: 10.1093/nar/25.17.3389
Alves, M., Pereira, A., Matos, P., Henriques, J., Vicente, C., Aikawa, T., et al. (2016). Bacterial community associated to the pine wilt disease insect vectors Monochamus galloprovincialis and Monachaons alternati. Sci. Rep. 6:23908. doi: 10.1038/srep23908
Alves, M., Pereira, A., Vicente, C., Matos, P., Henriques, J., Lopes, H., et al. (2018). The role of bacteria in pine wilt disease: insights from microbiome analysis. FEMS Microbiol. Ecol. 94:fiy077. doi: 10.1093/femsec/fiy077
Anke, M. K., Szymańska, K., Schwabe, R., Wiche, O., and Tischler, D. (2017). On the immobilization of desferrioxamine-like siderophores for selective metal binding. Solid State Phenom. 262, 517–520. doi: 10.4028/www.scientific.net/ssp.262.517
Arnow, L. E. (1937). Colorimetric estimation of the components of 3,4-dihydroxyphenylalanine-tyrosine mixtures. Comp. A J. Comp. Educ. 118, 531–537
Ashelford, K. E., Chuzhanova, N. A., Fry, J. C., Jones, A. J., and Weightman, A. J. (2006). New screening software that shows most recent large 16S rRNA gene clone libraries contain chimeras. Appl. Environ. Microbiol. 72, 5734–5741. doi: 10.1128/AEM.00556-056
Atkin, C. L., Neilands, J. B., and Phaff, H. J. (1970). Rhodotorulic acid from species of Leucosporidium, Rhodospiridium, Rhodotorula, Sporidiobolus, and Sporobolomyces, and a new alanine-containing ferrorhod from Cryptococcus melibiosus. J. Bacteriol. 103, 722–733
Aziz, R. K., Bartels, D., Best, A. A., DeJongh, M., Disz, T., Edwards, R. A., et al. (2008). The RAST Server: rapid annotations using subsystems technology. BMC Genomics 9:75. doi: 10.1186/1471-2164-9-75
Brenner, S. (1974). The genetics of Caenorhabditis elegans. Genetics 77, 71–94. doi: 10.1002/9780023003062
Chen, I.-M. A., Markowitz, V. M., Palaniappan, K., Szeto, E., Chu, K., Huang, J., et al. (2016). Supporting community annotation and user collaboration in the integrated microbial genomes (IMG) system. BMC Genomics 17:307. doi: 10.1186/s12864-016-2629-y
Choi, J. I., Yoon, K. H., Subbammal Kalichamy, S., Yoon, S. S., and Lee, J. II (2016). A natural odor attraction between lactic acid bacteria and the nematode Caenorhabditis elegans. ISME J. 10, 558–567. doi: 10.1038/ismej.2015.134
Dellagi, L., Segond, D., Rigault, M., Fagard, M., Simon, C., Saindrenan, P., et al. (2009). Microbial siderophores exert a subtle role in Arabidopsis during infection by manipulating the immune response and the iron status. Plant Physiol. 150, 1687–1696. doi: 10.1104/pp.109.138366
Esuola, C. O., Babalola, O. O., Heinie, T., Schwabe, R., Schlomann, M., and Tischler, D. (2016). Identification and characterization of a FAD-dependent putrescine N-hydroxylase (GorA) from Gordonia rubripertincta CW2B. J. Mol. Catal. B Enzym. 134, 378–389. doi: 10.1016/j.molcatb.2016.08.003
Fones, H., and Preston, G. M. (2013). The impact of transition metals on bacterial plant disease. FEMS Microbiol. Rev. 37, 495–519. doi: 10.1111/1574-6976.12004
Fonseca, L., Cardoso, J. M. S., Lopes, A., Pestana, M., Abreu, F., Nunes, N., and et al. (2012). The pinewood nematode, Bursaphelenchus xylophilus, in Madeira island. Helminthologia 49, 96–103. doi: 10.2478/s11867-012-0020-23
Gamalero, E., and Glick, B. R. (2011). “Mechanisms used by plant growth-promoting bacteria,” in Bacteria in Agrobiology: Plant Nutrient Management, ed. D. K. Maheshwari, (Berlin: Springer), 17–46. doi: 10.1007/978-3-642-21061-7
Gurevich, A., Saveliev, V., Vyakh, N., and Tesler, G. (2013). QUAST: quality assessment tool for genome assemblies. Bioinformatics 29, 1072–1075. doi: 10.1093/bioinformatics/btt086
Hider, R. C., and Kong, X. (2010). Chemistry and biology of siderophores. Nat. Prod. Rep. 27, 637–657. doi: 10.1039/904697a
Hirschmann, M., Grundmann, F., and Bode, B. H. (2017). Identification and occurrence of the hydroxamate siderophores aerobactin, putrebacint, avaroferrin and ochrobactin C as virulence factors from entomopathogenic bacteria. Environ. Microbiol. 19, 4080–4090. doi: 10.1111/1462-2920.13845
Huntemann, M., Ivanova, N. N., Movm rampatias, K., Tripp, H. J., Paez-Espino, D., Palaniappan, K., et al. (2015). The standard operating procedure of the DOE-JGI microbial genome annotation pipeline (MGAP v.4). Stand. Genomic Sci. 10, 86. doi: 10.1086/s04793-015-0077-y
Kang, D., Kirienko, D. R., Webster, P., Fisher, A. L., and Kirienko, N. V. (2018). A natural odor attraction between lactic acid bacteria and the nematode Caenorhabditis elegans, removes iron, and activates a distinct host response. Virulence 9, 804–817. doi: 10.1080/21505594.2018.1449508
Khan, M. R., Mohidin, F. A., Khan, U., and Ahamad, F. (2016). Native Pseudomonas spp. suppressed the root-knot nematode in in vitro and in vivo,
and promoted the nodulation and grain yield in the field grown mungbean. *Bioi. Control* 101, 159–168. doi: 10.1016/j.biocontrol.2016.06.012

Kim, O.-S., Cho, Y.-J., Lee, K., Yoon, S.-H., Kim, M., Na, H., et al. (2012). Introducing ExTaxon-e: a prokaryotic 16S rRNA gene sequence database with phylotypes that represent uncultured species. *Int. J. Syst. Evol. Microbiol.* 62, 716–721. doi: 10.1099/ijs.0.038075-38070

Kirienko, N. V., Kirienko, D. R., Larkins-Ford, J., Wählby, C., Ruvkun, G., and Ausubel, F. M. (2013). *Pseudomonas aeruginosa* disrupts *Caenorhabditis elegans* homeostasis, causing a hypoxic response and death. *Cell Host Microbe* 13, 406–416. doi: 10.1016/j.chom.2013.03.003

Kügler, J. H., Muhle-Goll, C., Hansen, S. H., Völp, A. R., Kirschhöfer, F., Kühl, B., Kim, O.-S., Cho, Y.-J., Lee, K., Yoon, S.-H., Kim, M., Na, H., et al. (2012). *Frontiers in Microbiology* | www.frontiersin.org

Proença, D. N., Francisco, R., Santos, C. V., Lopes, A., Fonseca, L., Abrantes, I. M. O., et al. (2010). Diversity of bacteria associated with *Rhodococcus erythropolis* B7g, a biosurfactant producing actinobacterium. *J. Biotechnol.* 280, 38–41. doi: 10.1016/j.jbiotec.2018.06.001

Ramaa, R., García, M. J., Lucena, C., Martínez-Medina, A., Aparicio, M. A., Ramos, J., et al. (2019). Induced systemic resistance (ISR) and fe deficiency responses in dicot plants. *Front. Plant Sci.* 10:287. doi: 10.3389/fpls.2019.00287

Saha, R., Saha, N., Donofrio, R. S., and Bestervelt, L. L. (2013). Microbial siderophores: a mini review. *J. Basic Microbiol.* 53, 303–317. doi: 10.1002/jobm.201100552

Saikia, R., Srivastava, A. K., Singh, K., Arora, D. K., and Lee, M.-W. (2005). Effect of iron availability on induction of systemic resistance to *Fusarium* wilt of Chickpea by *Pseudomonas spp.* *Mycolobiology* 33, 35–40. doi: 10.4489/MYCO.2005.33.1.035

Schneider, M., Tognoll, M., and Bairoch, A. (2004). The Swiss-Protein knowledgebase and ExPASy: providing the plant community with high quality proteomic data and tools. *Plant Physiol. Biochem.* 42, 1013–1021. doi: 10.1016/j.plaphy.2004.10.009

Senges, C. H. R., Al-Dilaimi, A., Marchbank, D. H., Wibberg, D., Winkler, A., Halit, B., et al. (2018). The secreted metabolome of *Streptomyces chartreusis* and implications for bacterial chemistry. *Proc. Natl. Acad. Sci. U. S. A.* 115, 2490–2495. doi: 10.1073/pnas.1751711115

Soutar, C. D., and Stavrinides, J. (2018). The evolution of three siderophore biosynthetic clusters in environmental and host-associating strains of *Pantoea*. *Mol. Genet. Genomics* 293, 1453–1467. doi: 10.1007/s00438-018-1477-7

Stamatakis, A. (2006). RAXML-VI-HPC: maximum likelihood-based phylogenetic analyses with thousands of taxa and mixed models. *Bioinformatics* 22, 2688–2690. doi: 10.1093/bioinformatics/btl446

Tash-Oshnoei, F., Harighi, B., and Abdollahzadeh, J. (2017). Isolation and identification of endophytic bacteria with plant growth promoting and biocontrol potential from oak trees. *Nat. Pathol.* 47, 1–8. doi: 10.1111/epf.12360

Vansuyt, G., Robin, A., Brat, J. F., Curie, C., and Lemanceau, P. (2007). Iron acquisition from Fe-pyoverdine by *Arabidopsis thaliana* extracellular proteases. *Bioinformatics* 23, 2688–2695. doi: 10.1093/bioinformatics/btm258

Vicente, C. S. L., Nascimento, F., Espada, M., Barbosa, P., Mota, M., Glick, B. R., et al. (2018). Microbial siderophores enterobactin promotes mitochondrial iron uptake and development of the host via interaction with ATP synthase. *Cell* 175:571–582.e111. doi: 10.1016/j.cell.2018.07.032

Wang, M., Carver, J. J., Phelan, V. A., Sanchez, L. M., Garg, N., Peng, Y., et al. (2016). Sharing and community curation of mass spectrometry data with global natural products social molecular networking. *Nat. Biotechnol.* 34, 828–837. doi: 10.1038/nbt.3597
Weber, T., Blin, K., Duddela, S., Krug, D., Kim, H. U., Bruccoleri, R., et al. (2015). antiSMASH 3.0—a comprehensive resource for the genome mining of biosynthetic gene clusters. *Nucleic Acids Res.* 43, W237–W243. doi: 10.1093/nar/gkv437

Wiche, O., Tischler, D., Fauser, C., Lodemann, J., and Heilmeier, H. (2017). Effects of citric acid and the siderophore desferrioxamine B (DFO-B) on the mobility of germanium and rare earth elements in soil and uptake in *Phalaris arundinacea*. *Int. J. Phytoremediation* 19, 746–754. doi: 10.1080/15226514.2017.1284752

Wilson, B. R., Bogdan, A. R., Miyazawa, M., Hashimoto, K., and Tsuji, Y. (2016). Siderophores in iron metabolism: from mechanism to therapy potential. *Trends Mol. Med.* 22, 1077–1090. doi: 10.1016/j.molmed.2016.10.005

Złoch, M., Thiem, D., Gadzała-Kopciuch, R., and Hrynkiewicz, K. (2016). Synthesis of siderophores by plant-associated metallotolerant bacteria under exposure to Cd2+. *Chemosphere* 156, 312–325. doi: 10.1016/j.chemosphere.2016.04.130

Zobel, R. W., Wright, S. F., Parker, D. R., Reichman, S. M., and Crowley, D. E. (2005). “Metal chelation in the rhizosphere,” in *Roots and soil management: interactions between roots and the soil*, eds R. W. Zobel, and S. F. Wright, (Madison, WI: American Society of Agronomy-Crop Science Society of America-Soil Science Society of America), 57–93.

**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2019 Proença, Heine, Senges, Bandow, Morais and Tischler. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.