Endophytic culturable bacteria colonizing *Lavandula dentata* L. plants: isolation, characterization and evaluation of their plant growth-promoting activities

Pereira, S.I.A.¹, Monteiro, C.¹, Vega, A.L.¹ and Castro, P.M.L.¹*

¹ CBQF – Centro de Biotecnologia e Química Fina – Laboratório Associado, Escola Superior de Biotecnologia, Universidade Católica Portuguesa/Porto, Rua Arquiteto Lobão Vital, Apartado 2511, 4202-401 Porto, Portugal

*Corresponding author

Paula M. L. Castro

CBQF - Centro de Biotecnologia e Química Fina – Laboratório Associado, Escola Superior de Biotecnologia, Universidade Católica Portuguesa/Porto

Rua Arquiteto Lobão Vital, Apartado 2511, 4202-401 Porto, Portugal

E-mail: plcastro@porto.ucp.pt

Tel: +351 22 558 0067

Fax: + 351 225 090351
Abstract

Aromatic plants such as lavender are stirring the attention of many researchers due to their content in bioactive secondary metabolites that can be used in traditional medicine. However, information regarding naturally occurring lavender associated bacterial endophytes (BE) is limited. To the best of our knowledge, this is the first study which aims to assess the phylogenetic diversity of the culturable endophytic bacteria of *Lavandula dentata* cultivated under organic management and to evaluate their potential as plant growth promoting (PGP) agents. BE were grouped by random amplified polymorphic DNA and identified by 16S ribosomal RNA gene sequencing. Endophytes were further characterized for the ability to produce several PGP substances, like ammonia, siderophores, indol-3-acetic acid, hydrogen cyanide and for the ability to solubilize phosphate. Plant cell-wall degrading enzymes were also determined. Densities of BE were higher in roots (log 6.39 CFU g⁻¹ fresh weight) than in shoots (log 5.56 CFU g⁻¹ fresh weight). Phylogenetic analysis showed that BE were affiliated to two major groups: γ-Proteobacteria (50%) and Firmicutes (31.6%) and a small part belonged to α- (7.9%) and β-Proteobacteria (10.5%), being *Pseudomonas* and *Bacillus* the most highly represented genera. Higher bacterial diversity was found in the lavender roots, with endophytes belonging to 6 different genera (*Pseudomonas, Variovorax, Rhizobium, Caulobacter, Bacillus* and *Paenibacillus*), than in shoots where only 3 genera (*Bacillus, Pseudomonas* and *Xanthomonas*) were found. Overall, BE showed ability to produce extracellular enzymes and multiple PGP traits, suggesting their potential use as efficient bioinoculants in sustainable cultivation of medicinal and aromatic plants.

Keywords

Endophytic bacteria; diversity; indol-3-acetic acid; lavender; plant growth promotion
1. Introduction

*Lavandula* species (Lamiaceae) are endemic to the Mediterranean region, Arabian Peninsula, Canary Islands, and India (Upson, 2002), being widely cultivated worldwide, due to their economic value for cosmetic and pharmaceutical industries and also for ornamental purpose (Bakkali et al., 2008; Muyima et al., 2002). Bioactive secondary metabolites such as essential oils are extracted from various aromatic plants and in particular those extracted from *Lavandula dentata* stand out for their use in popular medicine as antidiabetic, antihypertensive, and antiprotozoal agent (Al-Musayeib et al., 2012), explaining the growing interest of the pharmaceutical industries in these natural compounds.

Lavender species can be propagated by seeds or by woody stem cuttings. The propagation through seeds is usually slow, plants show a large variation in growth and in oil composition, and the genetic uniformity of the resulting plants is not guaranteed (Zuzarte et al., 2010). For these reasons, vegetative propagation is preferred to reproduction by seeds. Propagation by cuttings is fast, does not require special techniques and methods and is less expensive than *in vitro* micropropagation (Zuzarte et al., 2010). However, the poor rooting capacity of cuttings is often observed. In order to overcome this problem, plant growth regulators like the indole-3-butyric acid (IBA) and indole-3-acetic acid (IAA) are commonly used and showed to be efficient in increasing the adventitious root formation and the rooting capacity of cuttings (Bona et al., 2010; Kasem and El-baset, 2014). However, chemical application raises environmental concerns and increase production costs thus the use of sustainable practices in the production of medicinal and aromatic plants is imperative.

Plant growth promoting rhizobacteria (PGPR) have been used as bioinoculants for improving plant growth in a wide range of environments and crop species (Pereira and Castro, 2014a; Pereira et al., 2015). Several authors also reported that PGPR induced root formation in stem cuttings of several aromatic plants, namely in *Mentha piperita* (Kaymak et al., 2008).
and in *L. angustifolia* (Kasem and El-baset, 2014). However, very little is known about the effects of bacterial endophytes (BE) on the rooting capacity of cuttings.

Endophytic bacteria are defined as those that colonize the inner tissues of healthy plants without causing symptoms of disease or detrimental effect on their host (Schulz and Boyle, 2006). Similarly to rhizobacteria, BE promote plant growth by several mechanisms, including phosphate solubilization, nitrogen fixation, production of siderophores and ammonia and through the production of wide range of phytohormones, such as auxins, cytokinins and gibberellic acids (Pereira and Castro, 2014b; Verma et al., 2001). Endophytic bacteria have the advantage of proliferating within the plant tissue thus facing less competition for nutrients and being protected from high-stress environment of the soil (Sturz et al., 2000). Endophytes have been isolated from a wide range of plant species and tissues suggesting their ubiquitous existence in higher plants. However, only a few of all the plants existing on earth have ever been studied relative to their bacterial endophytic pool (Strobel and Daisy, 2003), increasing the probability to find new and beneficial endophytes with potential to be applied in biotechnology. The microbiome of medicinal and aromatic plants is extremely important since there is increasing evidences that the spectrum of bioactive metabolites may be related to the activity of associated BE (Emiliani et al., 2014). However, little attention has been paid to the BE from aromatic and medicinal plants and there is a lack of reports in this research area.

The aim of the present study was to isolate and characterize BE from tissues (root and shoot) of *L. dentata* plants growing under organic management. We further intended to assess the phylogenetic diversity and the phenotypic characterization of the culturable BE; and to evaluate their ability to produce plant growth promoting (PGP) substances, such as IAA, ammonia, hydrogen cyanide (HCN), siderophores and extracellular enzymes (lipases, cellulases, proteases, pectinases) and to solubilize phosphate, in order to indicate BE with...
potential to be used as bioinoculants in vegetative propagation of aromatic and medicinal plants.

2. Material and Methods

2.1 Enumeration and isolation of cultivable bacterial endophytes

To isolate representative BE four healthy *Lavandula dentata* L. plants grown under organic management were collected at random from the “Cantinho das Aromáticas” located in Vila Nova de Gaia, Portugal (GPS - 41°07’30.00”N; 8°38’40.82”W). Plant surface sterilization was performed according to Pereira and Castro (2014b). The success of the surface disinfection process was evaluated by plating 100 µl of water from the final rinse on Trypticase Soy Agar (TSA; Pronadisa) agar medium and incubated at 30 ºC for 3 days. No bacterial growth was found.

For bacterial enumeration 1 g of roots and shoots were homogenized in 9 ml of sterile phosphate buffered saline (PBS g l⁻¹: Na₂HPO₄, 1.44; KH₂PO₄, 0.24; KCl, 0.20; NaCl, 8.00; pH 7.4). Serial dilutions were made in duplicate and 0.1 ml of each dilution was spread onto TSA and Plate Count Agar (PCA; Merck) media. Plates were incubated for 7 days at 30 ºC and the number of colony forming units (CFU) determined. Different bacterial colonies were isolated based on size, morphology and color, from TSA and PCA media, using a streak-plate procedure. Isolates were purified and further phenotypically characterized.

2.2 Random Amplified Polymorphic DNA (RAPD) and DNA sequencing analysis

Bacterial isolates were grouped according to species similarity, based on RAPD profiles produced with primer M13 (5’GAGGGTGGCGGTTTCT-3’). DNA extraction and RAPD analysis were performed according to the methods described in Pereira and Castro (2014b). RAPD amplification products were run on a 1.5% agarose gel stained with SYBR
Safe (Invitrogen, UK) for 135 min at 80 V. RAPD patterns were compared using Bionumerics software (Applied Maths, St-Martens-Laten, Belgium) and clustered according to their similarities. Isolates displaying unique RAPD profiles were subsequently identified by 16S rRNA gene sequencing analysis. 16S rDNA amplification was performed with universal primers, 27F and 1492R as previously described by Pereira and Castro (2014b). The PCR products were purified using a GRS PCR & Gel Band Purification Kit (Grisp) and sequencing was performed by Macrogen Inc. (Netherlands). Sequence editing and inspection were performed using BioEdit program 7.0 version and the sequence similarity search was performed using the EzTaxon sever. For phylogenetic analyses, the sequences were aligned by using the CLUSTALW (Thompson et al., 1994). Tree constructions were performed with MEGA 5.0, using the neighbour-joining method (Kimura two-parameter distance optimized criteria). The robustness of the phylogenetic tree was confirmed by using bootstrap analysis based on 1000 resamplings of the sequences. The 16S rRNA sequences of the BE were deposited in GenBank database under accession numbers KP407086 to KP407123. Simpson’s (1-D) and Shannon’s (H) diversity indices were calculated based on the percentage of different bacterial genera in both plant compartments.

2.3 Plant growth promotion (PGP) traits

The amount of IAA produced by BE was determined according to Gordon and Weber (1951). Briefly, an aliquot of 500 μl of supernatant obtained from bacterial cultures grown in the presence of L-tryptophan (1 %) was mixed with 350 μl of Salper reagent. The absorbance of pink colour developed after 30 min of incubation in dark was read at 530 nm. The IAA concentration was determined using a calibration curve of pure IAA as a standard. Detection of siderophore production was carried out by inoculating the BE on Chrome Azurol S medium (Schwyn and Neilands, 1987). The development of a yellow to orange halo around
the colonies after incubation at 30 ºC for 72 h indicated a positive result for siderophore production. The detection of HCN production was made by amending nutrient agar with 4.4 g glycine l⁻¹ and streaking the isolates on this modified agar plates. On the top of each plate, a sterilized filter paper (Whatman No.1) soaked in 2% sodium carbonate prepared in 0.5 % picric acid solution was placed. Plates were incubated at 30 ºC for 4 days after which development of orange to red color indicated HCN production (Ahmad et al., 2008). For assessing the ability to produce ammonia, fresh cultures were inoculated into 10 ml of peptone water and incubated at 30 ºC for 48 h; following this, 0.5 ml of Nessler’s reagent (Sigma-Aldrich) were added to each tube and development of yellow to brown color was considered as a positive result for ammonia production (Cappuccino and Sherman, 1992). For phosphate solubilisation assay, fresh cultures were inoculated into National Botanical Research Institute Phosphate (NBRIP) medium supplemented with 0.5 % tricalcium phosphate (Nautiyal, 1999). The presence of a clearing halo around bacterial colonies were considered positive for phosphate solubilization.

Bacterial endophytes were screening for their ability to produce extracellular enzymes (proteases, cellulase, pectinase and lipases) according to the methods described in Pereira and Castro (2014b).

3. Results and Discussion

In the last years, several studies have focused their attention on aromatic and medicinal plants due to the presence of distinct bioactive secondary metabolites that can be used in traditional medicine. Despite the great interest in medicinal plants, very little is known about the associated BE. To the best of our knowledge, this work is the first to assess the phylogenetic diversity of the culturable endophytic bacterial populations of *L. dentata* plants cultivated under organic management and to evaluate their plant growth promoting traits.
In this study, the number of endophytic culturable bacteria varied significantly among plant organs. The colony forming units in roots varied between log 6.21 CFU g\(^{-1}\) fresh weight (FW) (for TSA medium) and log 6.39 CFU g\(^{-1}\) FW (for PCA medium) and were higher if compared to shoot where bacterial counts ranged from log 5.31 CFU g\(^{-1}\) FW (for TSA) to log 5.56 CFU g\(^{-1}\) FW (for PCA). Similar results were obtained for *Echinacea purpurea* and *Echinacea angustifolia* where the lowest bacterial counts were determined in the aboveground tissues, while the highest bacterial numbers were detected in roots (Chiellini et al., 2014). A similar trend was observed by Aravind et al. (2009) in black pepper (*Piper nigrum*) plants. However, several authors reported that population density of endophytes in aromatic and medicinal plants seem to be highest in aerial than in the belowground tissues. Emiliani et al. (2014) and El-Deeb et al. (2013) showed that leaves had higher numbers of endophytic bacteria than roots in *Lavandula angustifolia* and *Plectranthus tenuiflorus* plants, respectively.

Based on the distinct colony characteristics a total of 56 BE were isolated from the plant tissues of *L. dentata* plants growing under organic management. However, according to RAPD analysis only 38 different profiles were recognised, corresponding to 27 bacterial strains from roots (LR) and 11 strains from shoots (LS) (Table 1). The higher numbers of bacterial counts found in lavender roots, as well as the higher number of endophytes recovered from this tissue, may reflect the intimate contact of roots with soil rhizosphere, which facilitate the entry of bacteria into the root tissues (Kobayashi and Palumbo, 2000). In addition, the endophytic bacterial population densities reported in the present study were higher than the earlier reports in aromatic and medicinal plants such as *P. nigrum* (Aravind et al., 2009) and *L. angustifolia* (Emiliani et al., 2014). These results may be related to the management practices used for the cultivation of lavender plants, since organic management
seems to promote microbial activity in rhizosphere (Reilly et al., 2013), increasing the bacterial pool that can enter into plant tissues.

One isolate of each cluster was chosen for 16S rRNA partial gene sequence (Fig. 1). As shown in Table 1, all BE showed high similarities (98-100%) with their closest related species. Phylogenetic analyses based on 16S rRNA gene sequences showed that BE isolated from lavender tissues belonged to two main groups: \(\gamma\)-Proteobacteria (50%) and Firmicutes (31.6%), being a small portion affiliated to \(\alpha\)-Proteobacteria (7.9%) and \(\beta\)-Proteobacteria (10.5%). Overall, BE belonged to 7 different genera, being Pseudomonas (47%) and Bacillus (29%) the better represented in lavender plants (Table 1). A similar trend was already described by several authors for other aromatic and medicinal plants. Emiliani et al. (2014) reported that 51% of the BE isolated from \(L.\) angustifolia tissues belonged to Pseudomonas genus, while Vendan et al. (2010) showed a predominance of the genus Bacillus in ginseng plants. In fact, both genera have been identified as frequently occurring endophytes in several plant species (Hallmann and Berg, 2006).

The results also showed that BE colonized differently lavender tissues. In roots, where higher bacterial diversity was found, endophytes belonged to 6 different genera (Pseudomonas, Variovorax, Rhizobium, Caulobacter, Bacillus and Paenibacillus), while in shoots 91% of the endophytic isolates were affiliated to Bacillus and Pseudomonas and only one isolate belonged to Xanthomonas genus. Some of these genera have already been described in other aromatic and/or medicinal plants. Cho et al. (2007) isolated 63 different endophytic strains belonging to 13 different genera including Bacillus and Paenibacillus from the interior of ginseng tissues. Emiliani et al. (2014) also isolated endophytes affiliated to Rhizobium, Pseudomonas, Bacillus genera from \(L.\) angustifolia plants. However, to the best of our knowledge BE belonging to Caulobacter and Variovorax genera were never detected in lavender plant compartments. The differences among the culturable bacterial populations in
both lavender organs were also highlighted by the diversity indices calculated based on genera distribution, since Simpson’s and Shannon’s indices were higher in roots (D=0.74 and H=1.45, respectively) if compared to shoots (D=0.56 and H=0.86, respectively).

The distribution of BE in roots and shoots of *L. dentata* plants may be related to the distinct anatomical and nutritional conditions found in both plant compartments, leading to the creation of specific ecological niches for endophytic growth. According to Gaiero et al. (2013) endophyte distribution within plants depends on a combination of skills to colonize and the allocation of plant resources. In addition, according to Compańt et al. (2010) the secretion of cell-wall degrading enzymes by endophytes is an important trait for tissue colonization and bacterial spreading inside plants. In this study, the production of extracellular enzymes such as cellulases, pectinases, proteases and lipases by BE was assessed and the results are shown in Table 2. Bacterial endophytes had higher ability to produce pectinases (68%) and proteases (79%) than cellulases (50%) and lipases (58%). In general, endophytic bacteria from shoots showed higher ability to produce plant cell wall-degrading enzymes, especially pectinases and proteases, suggesting that endophytic colonization of shoots may be closely related to BE capacity to produce extracellular enzymes. Verma et al. (2001) also reported cellulase and pectinase activities in different isolates suggesting their potential for inter- and intracellular colonization.

Rooting of stem cutting is widely used in horticulture for the propagation of aromatic and medicinal plants, however poor rooting is often observed (Kasem and El-baset, 2014). In order to overcome this problem synthetic auxins are often applied at stimulating the adventitious root formation on cuttings (Bona et al., 2010), however due to the continuous increase of environmental footprint it is urgent to find eco-friendly alternatives. Although several studies have reported the beneficial effect of BE and PGPR inoculation in several
economically-important crops, like maize (Pereira and Castro, 2014a), carrot and potatoes (Surette et al., 2003), little information is available for aromatic and medicinal plants.

Screening results of PGP traits of the lavender BE are shown in Table 2. All endophytic strains were able to synthesize IAA in the presence of the precursor L-tryptophan, with levels ranging from 7.0 to 74.7 mg l\(^{-1}\). Several BE (21%) produced more than 40 mg l\(^{-1}\) of IAA and most of them were affiliated to \textit{Pseudomonas} genus. Bacterial endophytes isolated from ginseng plants also showed ability to synthetize IAA even though at lower levels (Vendan et al., 2010). The application of PGPR in order to minimize the use of synthetic auxins was already reported by Kasem and El-baset (2014) and Kaymak et al. (2008) who showed that the inoculation of PGPR in \textit{L. dentata} cuttings and \textit{M. piperina}, respectively improved rooting performance. However, so far no studies are available concerning the PGP of BE isolated from aromatic and medicinal plants and their ability to increase stem cutting establishment in soils. Additionally to auxin production, all endophytic strains were able to produce ammonia and siderophores. According to Idris et al. (2004) siderophore production among endophytes may be a general phenotype, since endophytes have to compete with plant cells for Fe supply. Several BE also showed ability to solubilize phosphate, which is of great importance to enhance P availability in soils during the initial colonization.

In this work, several potential human pathogens were isolated from lavender tissues, namely the strains LR 1-11, LR 2-1, LR 2-4, LR 2-6, LR 3-3, LR 3-7, LS 1-4 and LS 3-1 that are members of the \textit{Bacillus cereus} group of bacteria. These BE were found in both plant organs, but with higher incidence in roots. This is an issue of concern, since many genera including \textit{Burkholderia}, \textit{Enterobacter}, \textit{Bacillus} and \textit{Stenotrophomonas} have been identified as colonizers of the plant rhizosphere but some members can also successfully colonize human organs and tissues causing diseases (Berg et al., 2005). Consequently, the selection of
bacterial endophytes to be applied as bioinoculants in plant cultures should take in consideration the risk of pathogenicity to ensure that they do not inadvertently pose a threat to human health.

4. Conclusions

Bacterial endophytes seem to colonize differently lavender tissues. Beyond the higher numbers of BE found in the roots, phylogenetic analysis also showed clear differences between plant organs. In roots, a high diversity of genera was observed and BE were affiliated to *Pseudomonas*, *Variovorax*, *Rhizobium*, *Caulobacter*, *Bacillus* and *Paenibacillus*, while in shoots isolates belonged to *Bacillus*, *Pseudomonas* and *Xanthomonas* genera. The BE isolated in this study showed several plant growth promoting traits which suggest their potential for plant growth promotion. The use of such BE as efficient bioinoculants may constitute an interesting alternative to the application of chemical compounds, for sustainable cultivation of medicinal and aromatic plants.

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**Figure Captions**

**Fig. 1.** Neighbour-joining phylogenetic tree based on partial 16S rRNA sequences, showing the relationships between sequences of representative strains of bacterial endophytes and some of their closest phylogenetic relatives. Bootstrap values are shown at nodes. *Acidilobus saccharovorans* (NR115208.1) was used as outgroup. Bar indicates 0.05 substitutions per nucleotide position.
Table 1 – Phylogenetic affiliation to the closest relative and characterization of bacterial endophytes isolated from *Lavandula dentata* L.

<table>
<thead>
<tr>
<th>Strain</th>
<th>NCBI accession No.</th>
<th>Phylogenetic affiliation</th>
<th>Closest 16S rDNA match (accession number)</th>
<th>Similarity (%)</th>
<th>Gram reaction</th>
<th>Colony pigmentation</th>
<th>Cell morphology</th>
<th>Catalase</th>
<th>Oxidase</th>
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<td>LR 1-1</td>
<td>KP407086</td>
<td>γ-Proteobacteria</td>
<td><em>Pseudomonas brassicacearum</em> subsp. <em>neuroresistans</em> ATCC 49054 (EU391388)</td>
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<td>+</td>
<td>+</td>
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<td>+</td>
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<td>-</td>
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<td>Red</td>
<td>-</td>
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<td><em>Bacillus cereus</em> AGK 284 (AJ391844)</td>
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Table 2 - Characterization of bacterial endophytes for multiple plant growth promoting (PGP) traits: NH₃, siderophore, IAA, HCN and extracellular enzymes: cellulase, pectinase, protease and lipase production and phosphate solubilization ability.

IAA is expressed as means ± SE (n=5 to 12). (−) negative, (+) positive/weak, (++) intermediate, (+++) strong production.
Fig. 1

[Diagram showing bacterial distribution and species]

- **α-Proteobacteria** (73.5%)
  - LS 2-2 (KP407118)
  - LS 2-3 (KP407119)
  - LS 1-5 (KP407116)
  - LS 1-1 (KP407112)
  - Pseudomonas lutea (AY364537.1)
  - Pseudomonas graminis (Y11150.1)

- **β-Proteobacteria** (15.5%)
  - LR 1-9 (KP407091)
  - LR 3-2 (KP407105)
  - LR 3-6 (KP407107)
  - LS 3-5 (KP407122)
  - Pseudomonas fiscovaginiae (FJ483519.1)
  - Pseudomonas moorei (AM293666.1)
  - Pseudomonas jessenii (AF608259.1)

- **γ-Proteobacteria** (11.0%)
  - LR 1-5 (KP407088)
  - LR 2-3 (KP407098)
  - Pseudomonas punonensis (JQ344321.1)
  - LR 2-9 (KP407101)

- **Firmicutes** (5.0%)
  - LS 2-1 (KP407117)
  - Pseudomonas congelans (AJ492828.1)

- **Bacteroidetes**
  - LS 3-8 (KP407123)
  - Variorovorax soli (DQ432053.1)
  - Variorovorax boronicumulans (AB300597.1)
  - LR 1-3 (KP407094)
  - LR 1-6B (KP407089)
  - LR 2-10 (KP407102)

- **Caulobacter vibrioideis** (AJO09957.1)
  - LR 1-12 (KP407093)
  - Rhizobium sp. (FR570231.1)
  - LR 1-2 (KP407087)
  - Rhizobium grahamii (JF424608.1)
  - LR 3-11 (KP407112)

- **Pseudomonas kribbensis** (AF339123.1)
  - LR 2-11 (KP407103)

- **Bacillus aerophilus** (AJ831844.2)
  - LR 3-6 (KP407109)
  - LS 1-2 (KP407114)
  - Bacillus aryabhattii (EF114313.2)

- **Acidilobus saccharovorans** (NR115208.1)