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Evaluating *Eucalyptus* leaf colonization by *Brasilonema octagenarum* (Cyanobacteria, Scytonemataceae) using *in planta* experiments and genomics

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**ABSTRACT**

**Background.** *Brasilonema* is a cyanobacterial genus found on the surface of mineral substrates and plants such as bromeliads, orchids and eucalyptus. *B. octagenarum* stands out among cyanobacteria due to causing damage to the leaves of its host in an interaction not yet observed in other cyanobacteria. Previous studies revealed that *B. octagenarum* UFV-E1 is capable of leading eucalyptus leaves to suffer internal tissue damage and necrosis by unknown mechanisms. This work aimed to investigate the effects of *B. octagenarum* UFV-E1 inoculation on *Eucalyptus urograndis* and to uncover molecular mechanisms potentially involved in leaf damage by these cyanobacteria using a comparative genomics approach.

**Results.** Leaves from *E. urograndis* saplings were exposed for 30 days to *B. octagenarum* UFV-E1, which was followed by the characterization of its genome and its comparison with the genomes of four other *Brasilonema* strains isolated from phyllosphere and the surface of mineral substrates. While UFV-E1 inoculation caused an increase in root and stem dry mass of the host plants, the sites colonized by cyanobacteria on leaves presented a significant decrease in pigmentation, showing that the cyanobacterial mats have an effect on leaf cell structure. Genomic analyses revealed that all evaluated *Brasilonema* genomes harbored genes encoding molecules possibly involved in plant-pathogen interactions, such as hydrolases targeting plant cell walls and proteins similar to known virulence factors from plant pathogens. However, sequences related to the type III secretory system and effectors were not detected, suggesting that, even if any virulence factors could be expressed in contact with their hosts, they would not have the structural means to actively reach plant cytoplasm.

**Conclusions.** Leaf damage by this species is likely related to the blockage of access to sunlight by the efficient growth of cyanobacterial mats on the phyllosphere, which may hinder the photosynthetic machinery and prevent access to some essential molecules. These results reveal that the presence of cyanobacteria on leaf surfaces is not as universally beneficial as previously thought, since they may not merely provide the
products of nitrogen fixation to their hosts in exchange for physical support, but in some cases also hinder regular leaf physiology leading to tissue damage.

**Subjects** Bioinformatics, Genetics, Genomics, Microbiology, Plant Science

**Keywords** Cyanobacteriota, Oxyphotobacteria, Phyllosphere, Lignocellulolytic enzymes, Virulence factors, Horizontal gene transfer

**INTRODUCTION**

Cyanobacteria are microorganisms that colonize a wide variety of habitats, including the phyllosphere, either as free-living colonies or in symbiotic relationships. Cyanobacterial symbioses with eukaryotes are usually mutualistic, with cyanobionts providing partners with fixed carbon or nitrogen and/or defending them with toxins or sunscreens in exchange for protection from extreme environmental conditions and predation ([Usher, Bergman & Raven, 2007; Adams et al., 2013; Rikkinen, 2007](#)). *Brasilonema octagenarum* differs from other symbiotic cyanobacteria in this regard by presenting strains that are capable of damaging plant leaves ([Aguiar et al., 2008](#)), a behavior that as far as is currently known is unique in this phylum.

Thus far, *Brasilonema* strains have been found on subaerophytic habitats from tropical and subtropical environments of Brazil, French Antilles, Mexico, and USA with ten species described, namely *B. bromelae*, *B. angustatum*, *B. burkei*, *B. geniculatum*, *B. lichenoides*, *B. octagenarum*, *B. roberti-lamii*, *B. sennae*, *B. terrestris*, and *B. tolantongensis* ([Fiore et al., 2007; Aguiar et al., 2008; Sant’Anna et al., 2011; Vaccarino & Johansen, 2012; Becerra-Absalón et al., 2013; Rodarte et al., 2014; Miscoe et al., 2016; Villanueva et al., 2018; Villanueva et al., 2019](#)). The distribution of the genus *Brasilonema* is most likely pantropical ([Hauer, 2010; Kaštovský et al., 2010](#)), thus new species are likely to be found in previously unsurveyed geographical regions and habitats. Nevertheless, although other strains have been isolated from the phyllospheres of different host species, plant damage has so far only been observed in *B. octagenarum*.

Ultrastructural analyses have previously shown that *B. octagenarum* UFV-E1 mats can invade the mesophylls of *Eucalyptus* leaves under nursery conditions, which results in internal cell damage and necrosis and leads to reduced rates of photosynthesis and stomatal gas exchange, eventually causing reduction in growth and harming the productivity of the host plant ([Aguiar et al., 2008](#)). This cyanobacterium colonized the aerial parts of *Eucalyptus* saplings and mineral substrates at the vicinity of a greenhouse (Figs. 1A–1C), with leaves colonized by strain UFV-E1 showing damage to limbs, petiole, and apical buds (Figs. 1D–1H). The molecular mechanisms by which this interaction occurs have not been elucidated yet, and whether the cellular damage is caused directly by the cyanobacterium or indirectly by its epiphytic growth is also currently unknown.

Since similar leaf damage has not been previously observed in cyanobacteria-plant interactions, it is possible that *Brasilonema octagenarum* has acquired the capacity of infecting plants from phytopathogenic organisms by horizontal gene transfer. Therefore, this work aimed to experimentally reproduce under controlled conditions the effects of *B.*
Infection of *Eucalyptus urograndis* saplings by *Brasilonema octagenarum* under greenhouse conditions in Minas Gerais, southeast Brazil. Arrows point to cyanobacterial mats. (A) *E. urograndis* nurseries presenting visible growth of *B. octagenarum* mats on leaf surfaces. (B–C) *B. octagenarum* biofilms colonizing mineral substrates in contact with water drained from a greenhouse containing *E. urograndis* plants colonized by *B. octagenarum* UFV-E1. (D–H) Limbs, petiole, and apical buds of young *E. urograndis* leaves colonized by *B. octagenarum* UFV-E1 presenting pathogenic symptoms, including necrosis. Photo credit: Maione W. Franco.

octagenarum UFV-E1 colonization on eucalyptus leaves; to characterize its genome to search for molecular mechanisms potentially involved in leaf damage by this strain; and to evaluate whether horizontal gene transfer had any role on the evolution of these mechanisms. For these purposes, the aerial parts of *Eucalyptus urograndis* saplings were inoculated with *B. octagenarum* UFV-E1 cultures and subsequent changes in morphology, physiology and growth of the plants were evaluated. Afterwards, the genomes of *B. octagenarum* UFV-E1 and four other *Brasilonema* spp. strains were sequenced, characterized and compared and the presence of sequences orthologous to proteins previously reported as involved in plant-pathogen interactions was inquired.

**MATERIAL AND METHODS**

**Leaf colonization**

*Brasilonema octagenarum* was originally found and isolated by Aguiar et al. (2008) in an eucalypt nursery in Minas Gerais, southeastern region of Brazil. *B. octagenarum* UFV-E1 was grown in liquid BG-11 medium (Allen, 1968) at temperatures of 25 ± 2 °C, photoperiod...
of 16/8 h light/dark and irradiance of 115 \( \mu \)m photons m\(^{-2}\)s\(^{-1}\). After 30 days, the cultures were filtered using general purpose filter paper under sterile conditions and concentrated into 10% of the initial volume of 3 L to eliminate supernatants and facilitate the inoculation of leaves. Six *Eucalyptus urograndis* clonal plants were transplanted to 2 L pots containing substrate and acclimatized for two weeks in a greenhouse under the controlled conditions of 25–30 °C temperature and above 80% relative humidity. Cell suspensions were inoculated homogeneously on the aerial parts of each plant, except for six uninoculated plants used as controls. After 30 days, the cyanobacterial colonies were mechanically removed from leaf limbs using gentle fingertip movements and the middle portion of the leaves was used for extraction and quantification of pigment content. Three discs were taken from different, fresh leaves (Ø = 0.59 cm) in each control or inoculated plant and immersed in 7 mL of dimethylsulfoxide (DMSO) (Wellburn, 1994) in the dark for 24 h. Absorbance was determined at \( \lambda \) of 665.1, 680 and 450 for chlorophyll \( a \), chlorophyll \( b \) and carotenoids, respectively. Stem height and leaf area were evaluated, and stem, root and leaf dry masses were measured after oven dehydration at 75 °C for 72 h. The leaf area was measured using a leaf area integrator (Delta-T Devices, Cambridge, UK). The averages of the data obtained were compared with the Tukey test (significance level \( P < 0.05 \)) using SAS 9.2 (SAS Institute, Cary, USA).

### Anatomical evaluation

Leaf blade samples collected from inoculated and control plants were fixed with 50% FAA solution (formaldehyde, acetic acid and 70% ethanol) for 48 h and stored in 70% ethanol (Johansen, 1940). After the middle blade region was sectioned and the samples were embedded in Histosec (Merck, Kenilworth, USA), 10 \( \mu \)m cross sections were obtained with an automatic advance rotary microtome. Part of the cuts were stained with safranin/astra blue (Bukatsch, 1972) for observing the general structure and presence of phenolic compounds, and the other part was stained with scarlet Sudan (Sass, 1951) for cuticle observation. Next, the cuts were mounted on Permunt synthetic resin (Thermo Fisher Scientific, Waltham, USA). Sections were examined and images were captured with the Olympus AX-70 photomicroscope system (Olympus, Shinjuku, Japan) using the Image-Pro Discovery software (MediaCybernetics, Silver Spring, USA).

### High-throughput sequencing

*Brasilonema octagenarum* UFV-E1, *B. octagenarum* CENA114, *B. octagenarum* UFV-OR1, *B. bromeliae* SPC 951 and *Brasilonema* sp. UFV-L1 were grown as non-axenic, unicyanobacterial cultures in 125 mL Erlenmeyer flasks containing 50 mL of sterile BG-11\(_0\) liquid medium (Allen, 1968; Stanier et al., 1971). For the reduction of associated microbes, the samples were washed with ultrapure water, 0.05% Extran, 0.85% NaCl, and a solution composed of 50 mM NaCl, pH 7.5 10 mM Tris-HCl and pH 8.0 2.5 mM EDTA. After inoculation, incubation was carried out at 25 ± 1 °C with a photoperiod of 14 h of light and 10 h of darkness under fluorescent light of 40 \( \mu \)mol photons m\(^{-2}\) s\(^{-1}\). Two-week-old cultures were centrifuged for 10 min at 7,690 \( \times \) g and cells were collected for DNA extraction using a modified version of the protocol established by Lin et al. (2010).
Paired-ends libraries were produced for all strains with the Nextera XT DNA Sample Prep Kit (Illumina) according to manufacturer’s instructions. Additionally, mate-paired libraries were prepared from 8 kbp inserts for strains UFV-E1 and CENA114 using the Nextera Mate Pair Library Prep Kit (Illumina) according to the manufacturer’s protocols in order to improve assemblies and obtain complete genomes for the main strain of this study and the strain not isolated from phyllosphere. Sequencing was carried out in the HiSeq 2500 platform using the HiSeq v4 Reagent Kit (Illumina) following the instructions provided by the manufacturer.

**Genome assembly**

Bases with qualities lower than Phred 28, adapters and sequences shorter than 50 bp were removed from the datasets with Trimmomatic 0.36 (Bolger, Lohse & Usadel, 2014). Mate-paired libraries were further refined with NxTrim 0.4.2 (O’Connell et al., 2015) for the separation of true mate-paired reads. *De novo* assemblies were carried out using SPAdes 3.11.1 (Bankevich et al., 2012) and MaSuRCA 3.2.4 (Zimin et al., 2013). Kraken 1.0 (Wood & Salzberg, 2014) and MetaBAT 2.12.1 (Kang et al., 2015) were used for identifying cyanobacterial sequences among the assembled contigs. Platanus 1.2.4 (Kajitani et al., 2014) was used for further scaffolding and gap closing. Genome completeness and contamination were estimated with CheckM 1.0.7 (Parks et al., 2014).

**Genome annotation**

The assembled genomes were automatically annotated with the NCBI Prokaryotic Genome Annotation Pipeline (Tatusova et al., 2016). When necessary, manual curation of annotations was performed with Artemis 16.0.17 (Carver et al., 2012) and BLAST 2.6.0+ (Camacho et al., 2009). Functional identification of translated protein sequences into orthologous groups was performed with eggNOG 4.5.1 (Huerta-Cepas et al., 2016), Blast2GO 5.2 (Götz et al., 2008) and WEGO (Ye et al., 2018). The RAST server (Overbeek et al., 2014; Brettin et al., 2015) was used for predicting the subsystems present in the assembled genomes. Protein sequences from the Pathogen-Host Interactions Database version 4.5 (Urban et al., 2020) were used to uncover genes encoding proteins similar to known virulence factors by using TBLASTN with cut-off values of 90% for coverage and 60% for positive-scoring amino acid matches. Translated protein sequences for hypothetical proteins were retrieved from the assembled genomes and their domains were predicted with the Phobius web server (Käll, Krogh & Sonnhammer, 2007). Predictions for possible effectors from the type III secretion system were performed with the EffectiveDB web server (Eichinger et al., 2016).

**Prediction of mobile genetic elements**

The occurrence of mobile genetic elements in the genomes obtained was verified according to the protocols described in Alvarenga et al. (2018). Insertion sequences were predicted with OASIS 8/11/08 (Robinson, Lee & Marx, 2012), ISEScan 1.5.4.3 (Xie & Tang, 2017) and the ISfinder database (Siguier et al., 2006; Siguier et al., 2012). Prophages were predicted with PhiSpy 3.2 (Akhter, Aziz & Edwards, 2012) and VirSorter 1.0.3 (Roux, Enault & Hurwitz, 2015). CRISPRs were detected with MinCED 0.2.0.
For the uncovering of genomic regions with anomalous content, Alien_Hunter 1.7 (Vernikos & Parkhill, 2006), OligoWords 1.2.1 and SeqWord Sniffer 2.0 (Ganesan et al., 2008) were used.

**Genome comparisons and phylogenomic analyses**

OrthoANIu 1.2 (Yoon et al., 2017) and USEARCH 10.0.240 (Edgar, 2010) were used for estimating average nucleotide identity among the assembled genomes. Orthologous protein clusters shared among the strains were surveyed by the OrthoVenn server (Wang et al., 2015). The assembled genomes were included in a dataset with 154 complete and nearly complete cyanobacterial genomes retrieved from the NCBI RefSeq database (O’Leary et al., 2016) for selection of marker sequences with Metaxa 1.0.2 (Bengtsson et al., 2011) and Phyla-AMPHORA 03/19/13 (Wang & Wu, 2013). Multiple sequence alignments for sixty single-copy translated protein sequences shared by all genomes and 16S rRNA gene sequences were produced with MAFFT 7.309 (Katoh & Standley, 2013) and concatenated into a single alignment. A maximum likelihood phylogenomic tree was reconstructed from the concatenated translated proteins/16S rRNA gene alignment with RAxML 8.2.9 (Stamatakis, 2014) using the best-fit evolutionary models for each partition as calculated with the phylogenomics-tools pipeline (https://github.com/kbseah/phylogenomics-tools). The tree was visualized with FigTree 1.4.2 (http://tree.bio.ed.ac.uk/software/figtree/) and edited with Inkscape 0.92 (https://inkscape.org/).

**RESULTS**

*Eucalyptus* leaf inoculation

We first reinoculated *B. octagenarum* strain UFV-E1 on *E. urograndis* leaves to check if the cyanobacterial colonization and its effects previously found under natural conditions could be artificially replicated on seedlings. One month after inoculation, *E. urograndis* leaves colonized by strain UFV-E1 showed a significant decrease of 19% in the Chl a/Chl b ratio in the areas colonized by the cyanobacterium (Table 1). When the content of the pigments Chl a, Chl b and carotenes were analyzed individually, significant alteration in the colonized areas in relation to the control samples was not verified (Table 1). The inoculated plants had an increase in root and stem dry mass (19% and 23%, respectively) and stem height (7%), while leaf dry mass and area showed no significant differences between inoculated plants and controls (Table 2).

*B. octagenarum* UFV-E1 colonized *E. urograndis* blades at points scattered along leaf surfaces (Figs. 2A and 2B). After the removal of the cyanobacterial colonies (Fig. 2C), the regions previously covered by denser colonies showed alterations in pigmentation (Fig. 2D). Transverse sections of *E. urograndis* leaves from the control group showed a prominent central rib on the abaxial face, presenting a bicollateral vascular bundle on both sides and fundamental parenchyma (Fig. 3A). The epidermis was unstratified, covered by cuticle, showing stomata mainly on the abaxial face. The mesophyll was dorsinventral, composed of palisade parenchyma formed by one or two layers of cells, and lacunar parenchyma, formed by three or four layers (Fig. 3B). In the plants colonized by *B. octagenarum*, colonies were observed mainly on the adaxial face (Figs. 3C and 3D). No structural changes were
Table 1  Pigment content in leaf areas of *Eucalyptus urograndis* plants colonized by *Brasilonema octagenarum*. *Significant effect at $P \leq 0.05$, Tukey test.

| Sample          | Chl a (µg cm$^{-2}$) | Chl b (µg cm$^{-2}$) | Carotenoids (µg cm$^{-2}$) | Chl a/b* |
|-----------------|----------------------|-----------------------|----------------------------|-----------|
| controls        | 15.64 ± 0.47         | 6.78 ± 0.29           | 2.98 ± 0.21                 | 2.32 ± 0.10 |
| inoculated plants | 14.45 ± 0.44         | 7.42 ± 0.33           | 2.93 ± 0.30                 | 1.95 ± 0.06 |

Table 2  Growth variables evaluated in *E. urograndis* plants after inoculation with *B. octagenarum*. Significant effects at * $P \leq 0.05$ and ** $P \leq 0.01$, Tukey test.

| Sample          | Root dry weight (g) | Stem dry weight (g) | Leaves dry weight (g) | Leaf area (cm$^2$) | Stem height (cm) |
|-----------------|---------------------|---------------------|-----------------------|-------------------|-----------------|
| controls        | 15.11* ± 0.63       | 23.07** ± 1.01      | 21.22 ± 1.82          | 3,586.5 ± 255.3   | 83.3** ± 1.3    |
| inoculated plants | 17.94 ± 0.73       | 28.36 ± 0.99        | 22.47 ± 0.49          | 4,136.5 ± 147.5   | 89.5 ± 1.2     |

observed caused by colonization (Fig. 3C) in the regions of the central rib or the remainder of the leaf lamina (Fig. 3D). The cuticle remained without changes in the regions colonized by *B. octagenarum* (Figs. 3F and 3G) in comparison with the controls (Figs. 3E and 3G).

**Genomic sequencing and characterization**

The genome sequencing of *Brasilonema* strains resulted in the assembly of nearly complete genomes for *B. octagenarum* UFV-E1 and *B. octagenarum* CENA114 while draft genomes were obtained for *B. octagenarum* UFV-OR1, *B. bromeliae* SPC 951 and *Brasilonema* sp. UFV-L1 (Fig. 4A, Table 3). The genome sequences obtained in this work were deposited in the NCBI Genome database under the accession numbers CP030118–CP030123, QMEA00000000, QMEB00000000, QMEC00000000. Predictions for the nearly complete chromosomes of strains UFV-E1 and CENA114 uncovered 17 and 14 genome islands, respectively. Additionally, 158 insertion sequences were found in both sequences. No prophages were detected in those chromosomes. On the other hand, two remnant prophage regions were predicted in each plasmid from both strains. Except for the UFV-E1 190 kbp plasmid, two genome islands were also detected in those plasmids. Moreover, 11 additional insertion sequences were found in the 250 kbp plasmids, while 18 and 16 insertion sequences were detected in the 190 kbp plasmids from UFV-E1 and CENA114, respectively. Finally, 9 CRISPR loci were predicted in both nearly complete chromosomes. Average nucleotide identity (ANI) among strains UFV-E1, CENA114, and UFV-OR1 was calculated as 99%, showing that they are actually representatives of the same species, *B. octagenarum*, which presented 90 and 85 ANI percentages when their genomes were compared to the genomes of *B. bromeliae* SPC 951 and *Brasilonema* sp. UFV-L1, respectively (Fig. 4B). Clustering analyses revealed a large number of orthologous protein clusters shared among the analyzed *Brasilonema* genomes, with a very small amount of protein clusters exclusive to each strain (Fig. 4C).

In the virulence, disease and defense subsystem predicted in the nearly complete *B. octagenarum* genomes, genes involved in resistance to fluoroquinolone- and betalactam-based antibiotics were found, in addition to genes for resistance to copper, mercury, arsenic,
Figure 2  *Eucalyptus urograndis* leaves colonized by *Brasilonema octagenarum* UFV-E1. (A) General appearance of colonized leaves presenting conspicuous growth of cyanobacterial filaments. (B) Cyanobacterial mats covering the central vein of the host leaf. (C) Removal of cyanobacterial colonies from the leaf surface. (D) Discoloration in leaf areas previously covered by cyanobacterial colonies. Photo credit: Maione W. Franco.

Chromium and cobalt/zinc/cadmium, suggesting that these cyanobacteria may have some tolerance to environmental disturbances, such as contamination by toxic element. However, subsystem coverage was very low in the assembled genomes, as merely 18% of sequences in the *B. octagenarum* genomes could be included into known subsystems, while subsystem coverage in the *B. bromeliaceae* SPC 951 and *Brasilonema* sp. UFV-L1 genomes amounted to 17% of their content. Similarly, the most abundant functional categories for clusters of orthologous groups (COGs) in all genomes were those with general function prediction only or unknown function. Overall all strains presented very similar number of genes
related to known COG and GO terms (Fig. 5, Table S2), suggesting that these species may perform similar ecological roles. The analysis of translated protein sequences from genes annotated as encoding hypothetical proteins with the Phobius webserver revealed that roughly three quarters are likely non-cytoplasmic and approximately one quarter of these
sequences present transmembrane domains (Table 4). This suggests that a considerable number of yet uncharacterized proteins are exported from the cells, with some of them possibly involved in ecological interactions.

In accordance with the latest system of cyanobacterial classification by Komárek et al. (2014), the phylogenomic analysis showed that the Brasilonema strains were closely related to Scytonema representatives, thus confirming previous polyphasic descriptions of the family Scytonemataceae, which in turn clustered inside the order Nostocales (Fig. 6), as expected based on previous works on the taxonomic positioning of these cyanobacteria (Fiore et al., 2007; Komárek et al., 2014).

Evaluation of sequences that might be involved in plant interactions

Twenty five secondary metabolite gene clusters were predicted in the nearly complete genomes of strains UFV-E1 and CENA114, including clusters encoding the pathways for polyketides, non-ribosomal peptides, post-translationally-modified peptides, bacteriocins, terpenes, indoles, and phenazines, most of which presented low similarity to reference clusters involved in the biosynthesis of known molecules. Five gene clusters, however, presented 100% similarity to references for clusters that encode the biosynthesis of the cyclic non-ribosomal peptides anabaenopeptin and nostopeptolide, the sesquiterpene geosmin, the polyketide 1-heptadecene, and the UV-protecting amino acid shinorine.
Table 3  Genomic features of the *Brasilonema* spp. strains characterized in this work.

| Feature                      | UFV-E1   | CENA114  | UFV-OR1  | SPC 951  | UFV-L1  |
|------------------------------|----------|----------|----------|----------|----------|
| completeness (%)             | 99.52    | 100.00   | 99.28    | 90.48    | 99.40    |
| contamination (%)            | 2.89     | 2.17     | 3.01     | 2.37     | 1.20     |
| total size (bp)              | 8,258,380| 8,226,398| 8,094,337| 6,277,767| 8,321,598|
| −chromosome (bp)             | 7,816,330| 7,782,641| —        | —        | —        |
| −plasmid 1 (bp)              | 251,877  | 252,036  | —        | —        | —        |
| −plasmid 2 (bp)              | 190,173  | 191,721  | —        | —        | —        |
| coding sequences (bp)        | 6,559,505| 6,529,766| 6,271,238| 4,742,865| 6,443,634|
| −chromosome (bp)             | 6,198,811| 6,170,908| —        | —        | —        |
| −plasmid 1 (bp)              | 204,544  | 204,193  | —        | —        | —        |
| −plasmid 2 (bp)              | 156,150  | 154,665  | —        | —        | —        |
| GC content (%)               | 42.49    | 42.50    | 42.48    | 42.97    | 41.97    |
| protein-coding genes         | 6,395    | 6,371    | 6,309    | 5,207    | 6,652    |
| pseudogenes                  | 497      | 502      | 541      | 432      | 516      |
| RNA genes                    | 58       | 56       | 58       | 54       | 46       |
| ncRNA                        | 4        | 4        | 4        | 4        |
| genes with assigned subsystems| 1,394    | 1,388    | 1,427    | 1,137    | 1,397    |
| genes assigned to COG categories with known function | 3,440 | 3,426 | 3,379 | 2,741 | 3,389 |
| hypothetical proteins        | 2,081    | 2,092    | 2,074    | 1,695    | 2,314    |

Similarity to references for the biosynthesis of known cyanotoxins, which might be involved in plant-pathogen interactions, was not found in any of the assembled genomes.

Annotations in all the *Brasilonema* genomes uncovered several genes encoding enzymes that likely have cellulose, lignin and xylan as substrates (Table 5), which may therefore have a role on leaf colonization and interact with host cell walls. Other enzymes that might have some role during plant colonization have also been annotated in the genomes of all the *Brasilonema* strains, including proteins involved in chitin and protein degradation, auxin biosynthesis and biofilm formation through exopolysaccharide biosynthesis (Table 6).

Protein sequences with similarity to relevant virulence factors from plant pathogens deposited in the Pathogen Host Interaction database were found encoded in the genomes of all five *Brasilonema* strains analyzed in the present work (Table 7). Similarities were found with proteins from phytopathogenic bacteria (*Burkholderia glumae*, *Erwinia amylovora*, *Pantoea ananatis*, *Ralstonia solanacearum*, *Xanthomonas campestris*, *X. citri* and *X. oryzae*), fungi (*Botrytis cinerea*, *Colletotrichum lagenaria*, *Fusarium graminearum*, *F. oxysporum*, *Magnaporthe oryzae* and *Parastagonospora nodorum*) and nematodes (*Bursaphelenchus xylophilus*), which target leaves, shoots, roots, cones, panicles, seedlings, cotyledons or fruits. Among the set of virulence factors with homologous sequences in the *Brasilonema* genomes (36 proteins), the majority (26 proteins) can be considered essential to phytopathogenicity.
Figure 5   Functional annotations of genes included in the *Brasilonema* spp. genomes characterized in this work, evidencing high similarity between strains and species in this genus. (A) Gene Ontology terms associated with the predicted properties of the products encoded in *Brasilonema* genomes. (B) Distribution of COG categories among *Brasilonema* genomes, showing that they encode a large number of poorly characterized proteins. COG categories W, Y, and Z did not present any matches and thus were suppressed.

Since the virulence of the organisms that express these factors may be reduced or even lost after deleterious mutations occur in their coding sequences (Table 7).

It is noteworthy that both nearly complete *B. octagenarum* genomes presented regions with content showing divergent composition when compared to their average content, which is commonly found in horizontally-acquired sequences, while further evidences of horizontal transfer were obtained by predictions of other mobile genetic elements (Fig. 4A).
Figure 6  Phylogenomic tree reconstructed by maximum likelihood. Analysis based on alignments of sequences of the 16S rRNA gene and sixty proteins codified in single-copy genes present in the genomes from the strains evaluated in the present work (highlighted in bold) and nearly complete cyanobacterial genomes and chromosome scaffolds from the NCBI RefSeq database.

Table 4  Domains predicted from hypothetical protein sequences encoded in Brasilonema genomes.

| Strain                        | Cytoplasmic | Non cytoplasmic | Non cytoplasmic + signal peptide | Cytoplasmic + non cytoplasmic + transmembrane | Cytoplasmic + non cytoplasmic + transmembrane + signal peptide |
|------------------------------|-------------|-----------------|---------------------------------|-----------------------------------------------|---------------------------------------------------------------|
| B. octogenarum UFV-E1        | 98          | 1,200           | 324                             | 417                                           | 42                                                            |
| B. octogenarum CENA114       | 96          | 1,206           | 329                             | 415                                           | 44                                                            |
| B. octogenarum UFV-OR1       | 99          | 1,195           | 313                             | 399                                           | 42                                                            |
| B. bromeliae SPC 951         | 83          | 1,019           | 204                             | 316                                           | 44                                                            |
| Brasilonema sp. UFV-L1       | 140         | 1,410           | 278                             | 416                                           | 50                                                            |

However, none of the sequences similar to known virulence factors in the Brasilonema genomes were found within regions estimated as originating from horizontal transfer, and thus these genes are unlikely to be of xenologous origin.
Table 5  
Annotations for enzymes encoded in Brasilonema genomes potentially having plant polysaccharides as substrates.

| Protein                               | Substrate                  | UFV-E1 | CENA114 | UFV-OR1 | SPC 951 | UFV-L1 |
|---------------------------------------|----------------------------|--------|---------|---------|---------|--------|
| cellulase M                           | cellulose                  | +      | +       | +       | +       | +      |
| 1,4-beta-glucanase / beta-1,4-glucanase| cellulose                  | +      | +       | +       | +       | +      |
| endo-1,4-beta-glucanase               | cellulose                  | +      | +       | +       | +       | +      |
| endo-1,4-beta-glucanase E1 precursor  | cellulose                  | +      | +       | +       | +       | +      |
| endo-1,4-beta-xylanase                | xylan                      | +      | +       | +       | +       | +      |
| endo-1,4-beta-xylanase Z precursor    | xylan                      | +      | +       | +       | +       | +      |
| alpha-glucosidase                    | glucoside                  | +      | +       | +       | +       | +      |
| beta-glucosidase / beta-fucosidase    | glucoside / fucoside       | +      | +       | +       | –       | –      |
| 6-fosfo-beta-glucosidase / beta-galactosidase | glucoside / galactoside   | +      | +       | +       | –       | +      |
| malto-oligosyltrehalose trehalohydrolase | malto-oligosyltrehalose   | +      | +       | +       | –       | –      |
| 4-carboxymuconolactone                | lignin                     | +      | +       | +       | –       | +      |
| decarboxylase/alkylhydroperoxidase    | lignin                     | –      | –       | –       | +       | +      |
| catalase KatE                         | lignin                     | –      | –       | –       | –       | +      |
| catalase-peroxidase KatG              | lignin                     | +      | +       | +       | –       | –      |
| catalase-like hemeprotein             | lignin                     | +      | +       | +       | +       | +      |
| manganese catalase                    | lignin                     | +      | +       | +       | +       | +      |
| dye-decoloring peroxidase             | lignin                     | +      | +       | +       | –       | –      |
| Bcp-type thiol peroxidase             | lignin                     | +      | +       | +       | +       | +      |
| alpha-amylase / alpha-manosidase      | starch                     | +      | +       | +       | +       | +      |

DISCUSSION

The initial growth of B. octagenarum on the surface of E. urograndis leaves apparently occurred as a relationship of epiphytism, as no detrimental effects to the host plant were initially observed. B. octagenarum colonization of E. urograndis later provided benefits to the host, which was evidenced by the higher production of root and stem dry mass in comparison to the control group (Table 2). The production and release of plant growth regulators by cyanobacteria has been confirmed in several studies with the identification of abscisic acid, auxins, cytokinins, ethylene and gibberelins (Singh et al., 2017), and stimulus to plant growth caused by cyanobacteria has been observed when these microorganisms are used as biofertilizers (Jhala, Panpatte & Vyas, 2017; Muñoz Rojas et al., 2018). Therefore, it is likely that colonization by cyanobacteria stimulated the growth of the host plants by the release of plant growth regulators by B. octagenarum.

On the other hand, the consequential decrease in the Chl a/b ratio in leaves resulting in the lighter green areas observed in Fig. 2 is most likely a direct response to the decrease in light incidence caused by sunlight blockage by the epiphyllic cyanobacterial colonies. Chl a/b ratio tends to decrease in response to shading, as previously observed in experiments with decreased light incidence in Tetrastigma hemsleyanum (Dai et al., 2009). This likely represents the early stages of leaf damage by the cyanobacterium and could lead to more severe consequences to the host plant in case of prolonged exposure. As described by Aguiar et al. (2008), in plants that remain for long periods under nursery conditions favorable to the development of microbial mats, extended cyanobacterial colonization starts to act as a
Table 6  Proteins encoded in the genomes of *Brasilonema* spp. strains presenting functional predictions previously described as involved in plant-infection processes.

| Protein                                      | Function                        | UFV-E1 | CENA114 | UFV-OR1 | SPC 951 | UFV-L1 |
|----------------------------------------------|---------------------------------|--------|---------|---------|---------|--------|
| glucosamine-6-phosphate deaminase            | chitin degradation              | +      | +       | +       | +       | +      |
| N-acetylglucosamine-6-phosphate deacetylase  | chitin degradation              | +      | +       | +       | +       | +      |
| N-acetylglucosamine kinase                   | chitin degradation              | +      | +       | +       | −       | +      |
| N-acetylglucosamine-related transporter NagX  | chitin degradation              | +      | +       | +       | +       | +      |
| ClpP class periplasmic serine protease       | protein degradation             | +      | +       | +       | +       | +      |
| serine protease                              | protein degradation             | +      | +       | +       | +       | +      |
| rhomboid family serine protease              | protein degradation             | +      | +       | +       | +       | +      |
| serpin family serine protease inhibitor      | protein degradation             | +      | +       | +       | +       | +      |
| anthranilate phosphoribosyltransferase       | auxin biosynthesis              | +      | +       | +       | +       | +      |
| phosphoribosylanthranilate isomerase         | auxin biosynthesis              | +      | +       | +       | +       | +      |
| aromatic L-amino acid decarboxylase          | auxin biosynthesis              | +      | +       | +       | −       | +      |
| indole-3-pyruvate decarboxylase              | auxin biosynthesis              | +      | +       | +       | −       | −      |
| tryptophan synthase –alpha chain             | auxin biosynthesis              | +      | +       | +       | +       | +      |
| tryptophan synthase –beta chain              | auxin biosynthesis              | +      | +       | +       | +       | +      |
| auxin efflux carrier protein                 | auxin biosynthesis              | +      | +       | +       | +       | +      |
| exopolysaccharide synthesis protein          | exopolysaccharide biosynthesis  | +      | +       | +       | −       | +      |
| exopolysaccharide production protein ExoZ     | exopolysaccharide biosynthesis  | +      | +       | +       | +       | +      |
| exopolysaccharide production protein ExoQ     | exopolysaccharide biosynthesis  | +      | +       | +       | −       | +      |
| Hop protein                                  | type III secretion system effector | +    | +       | +       | +       | +      |

stress factor compromising the source/drain balance of eucalyptus plants and could result in necrosis.

Overall, the *Brasilonema* protein sequences that presented similarity to sequences in PHI-base could perform roles in diverse molecular mechanisms such as biosynthesis and degradation, transport, signaling, regulation, starvation response, reproduction, motility and other processes, including currently unknown functions. Ten matches between the translated protein sequences from the *Brasilonema* genomes and proteins deposited in PHI-base (C5A9K4, D4I307, F5HCK8, G4ML75, G4MTK2, Q4UTV7, Q4UUL4, Q8XSV8, Q8Y0J2 and Q9C1T0) are likely to indicate enzymes with roles in primary metabolism (Cooley et al., 1999; Namiki et al., 2007; O’Connell et al., 2013; Karki & Ham, 2014; Ramos et al., 2014; Feng et al., 2015; Saint-Macary et al., 2015; Zhang et al., 2015a). Since mutants for the inactivation of the genes that encode their homologues in pathogens usually present auxotrophy, the presence of genes for proteins homologous to those factors in the *Brasilonema* genomes does not necessarily reflect any additional capacity of these strains to colonize plant surfaces. Additionally, a number of sequences in the *Brasilonema* genomes have presented similarity with polyketide synthases of unknown functions deposited in PHI-base (UniProt accession numbers I1RSU2, Q6RKH3, Q6RKH4 and Q6RKH5), but since the inactivation of the genes encoding these enzymes have not affected pathogenicity, it is unlikely that their homologues in *Brasilonema* genomes have roles in plant colonization. Four sequences from the UFV-E1 genome matched proteins that have been described as part of two-component systems involved in pathogenic
### Table 7  Virulence factors synthesized by phytopathogens with similarity to proteins encoded in *Brasilonema* genomes. An expanded version of this table is provided as Table S1.

| UniProt accession | Protein       | Function                                      | UFV-E1 | CENA114 | UFV-OR1 | SPC 951 | UFV-L1 |
|-------------------|---------------|-----------------------------------------------|--------|---------|---------|---------|--------|
| AOA0K0GHK1        | SreR          | two-component system regulation               | +      | +       | +       | +       | +      |
| AOA0K0GI30        | DetR          | two-component system cytoplasmic signaling    | +      | +       | +       | +       | +      |
| A0A0U1YU79        | PiIT          | twitching motility                            | +      | +       | +       | +       | +      |
| A4K9H16           | PhoP          | two-component system regulation               | +      | +       | +       | +       | +      |
| A4QUT2            | CPXB          | catalase-peroxidase                           | +      | +       | +       | +       | +      |
| B0LFQ7            | Bx-Prx        | reproduction and pathogenicity                | +      | +       | +       | +       | +      |
| C5A846            | PidR          | two-component system regulation               | +      | +       | +       | +       | +      |
| C5A9K4            | AroB          | 3-dehydroquinate synthase                     | +      | +       | +       | –       | +      |
| D4HUW4            | YhbH          | ribosome-associated modulation protein        | +      | +       | +       | +       | +      |
| D4HX24            | RpoS          | regulation of stress and starvation response  | –      | –       | –       | +       | +      |
| D4HXR8            | AcrD          | resistance-nodulation-cell division transport | +      | +       | +       | +       | +      |
| D4H307            | ArgD          | N-acetylornithine aminotransferase            | +      | +       | –       | –       | +      |
| F5HCK8            | IPMDH         | 3-isopropylmalate dehydrogenase               | +      | +       | +       | +       | +      |
| G4ML75            | MET6          | methylnitrohydroxypteroylglutamate-homocysteine S-methyltransferase | +      | +       | –       | –       | +      |
| G4MTK2            | MoARG1        | arginine biosynthesis                         | +      | +       | +       | +       | +      |
| G4MXC5            | MoPRX1        | peroxiredoxin peroxidase                      | +      | +       | +       | +       | +      |
| G4N4N6            | MoSFA1        | S-(hydroxymethyl)glutathione dehydrogenase    | +      | +       | +       | +       | +      |
| G4NCL5            | MGG_00383     | hypothetical protein                          | +      | +       | +       | +       | +      |
| I1RSU2            | PKS9/KSA1     | polyketide synthase                           | +      | +       | +       | +       | –      |
| P87025            | THR1          | reductase                                     | –      | –       | –       | +       | +      |
| Q12634            | BUF/BUF1      | undefined                                     | +      | +       | +       | +       | +      |
| Q2L9K2            | BcPICS/BcFKB12 | rapamycin sensitivity                         | +      | +       | +       | +       | +      |
| Q4UQDO            | XC_3703       | cyclic di-GMP effector                        | +      | +       | +       | –       | +      |
| Q4UTV7            | XC_2466       | aspartate alpha-decarboxylase                 | +      | +       | +       | +       | +      |
| Q4UUU4            | XC_2203       | nucleotide diphosphate kinase                 | +      | +       | +       | +       | +      |
| Q58PW8            | HsvA          | hrp-associated systemic virulence             | –      | –       | –       | –       | +      |
| Q5H3K9            | ColR          | undefined                                     | +      | +       | +       | +       | +      |
| Q6RKH3            | PKS7          | polyketide synthase                           | +      | +       | +       | –       | +      |
| Q6RKH4            | PKS6          | polyketide synthase                           | +      | +       | +       | –       | +      |
| Q6RKH5            | PKS5          | polyketide synthase                           | +      | +       | +       | –       | –      |
| Q7WTQ9            | AcrB          | multidrug efflux pump                         | +      | +       | +       | +       | +      |
| Q8PM59            | PstB          | phosphate metabolism                          | +      | +       | +       | +       | +      |
| Q8XSV8            | FabG2         | 3-ketoacyl-acyl carrier protein reductase      | +      | +       | +       | +       | +      |
| Q8YOJ2            | FabG1         | 3-ketoacyl-acyl carrier protein reductase      | +      | +       | +       | +       | +      |

(continued on next page)
processes (accessions A0A0K0GHK1, A0A0K0GI30, A4K9H6 and C5A846), but since two-component systems control behavior not only during pathogenicity, but also in cell communication and environmental adaptation (Zschiedrich, Keidel & Szurmant, 2016), these proteins might also have biological roles other than plant colonization.

The remaining protein sequences with similarity to PHI-base entries could have important roles for plant colonization. Five Brasilonema sequences have presented similarity with proteins involved in regulatory cascades for virulence (D4HUY4, D4HX24, Q4UQD0, Q58PW8 and Q5H3K9) (Oh, Kim & Beer, 2005; Subramoni et al., 2012; An et al., 2014; Ancona, Li & Zhao, 2014; Santander et al., 2014). Two hypothetical proteins from PHI-base (G4NCL5 and Q2LK92) presenting similarity to Brasilonema sequences were previously shown to have significant effects on plant pathogenicity, including severe symptoms like necrosis (Gioti et al., 2006; Jeon et al., 2007). Three copies of the twitching motility gene pilT were annotated in the UFV-E1 genome, important for spreading pathogen cells on the surface of the host (Weller-Stuart et al., 2017). Three translated protein sequences from the UFV-E1 genome have shown similarity with catalase-peroxidases and peroxiredoxins from PHI-base (accessions A4QUT2, B0LFQ7 and G4MXC5), enzymes with very important roles in the maintenance of redox homeostasis and the protection of pathogen cells from oxidative damage brought by hydrogen peroxide produced by host cells (Tanabe et al., 2011; Fu et al., 2014; Mir et al., 2015; Santander, Figàs-Segura & Biosca, 2018). Genes for other proteins potentially involved in detoxification have also been predicted, including multidrug efflux pumps (D4HX8 and Q7WTQ9), which may confer resistance to phytoalexins (Burse, Weingart & Ullrich, 2004), and S-(hydroxymethyl)gluthatione dehydrogenase (G4N4N6), which is produced by phytopathogens as protection against damage from nitric oxide (Zhang et al., 2015b).

Although the identification of sequences similar to virulence factors in the Brasilonema genomes represents a potential for these cyanobacteria to cause damage to plants, their role within Brasilonema interactions with their hosts warrants further functional studies for the evaluation of the conditions under which these genes can be expressed. Nevertheless, except for one gene encoding a putative secreted protein that is likely a distant homolog of the Hop effector protein (Table 5), which is also found in other cyanobacteria with a presumed function of kinase (Zhang et al., 2016), the type III secretion system as well as its effectors appear to be completely absent in the assembled Brasilonema genomes. This is in agreement with observations for other cyanobacteria showing that they present the types I, II and IV secretion systems, but not the type III (Gonçalves et al., 2019). Since the transmission of this secretion system can occur either by vertical or horizontal transfer (Jackson et al., 2011), its absence in Brasilonema spp. together with the lack of evidence for horizontal transfer in the encoding of other molecules provides further evidence that

### Table 7 (continued)

| UniProt accession | Protein | Function                           | UFV-E1 | CENA114 | UFV-OR1 | SPC 951 | UFV-L1 |
|--------------------|---------|------------------------------------|--------|---------|---------|---------|--------|
| Q9C1T0             | ARG1    | argininosuccinate lyase             | +      | +       | +       | +       | +      |
| Q9UWF0             | CAM     | undefined                          | –      | –       | –       | –       | +      |

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leaf damage in this species emerged independently and not from the acquisition of genetic material from plant pathogens.

The role of type III secretion systems together with cell wall-degrading enzymes, serine proteases, phytohormones, siderophores and exopolysaccharides, among other molecules, has been increasingly recognized in plant pathogenesis (Lindeberg, 2012; Melotto & Kunkel, 2013). Most phytopathogens that affect plant productivity (including species of Xanthomonas, Pseudomonas, Agrobacterium, Ralstonia and Erwinia) are gram-negative, as are cyanobacteria, and one of the main mechanisms of virulence and pathogenicity of these phytopathogens is the type III secretion system, through which effector proteins are injected into host tissues, bringing about several functions that promote pathogenic processes (Sherif et al., 2015). Consequently, even if potential virulence factors, plant cell wall-degrading enzymes and other molecules were produced by the cyanobacteria in contact with their hosts, the absence of the type III secretion system in Brasilonema spp. is therefore a great obstacle for these microbes to actively infect plant cells.

CONCLUSIONS

Brasilonema octagenarum UFV-E1 efficient growth and colonization of the phyllosphere was shown to cause lighter green areas on Eucalyptus urograndis leaves as a result of the decrease in the Chl a/b ratio. Sequences similar to known virulence factors from phytopathogens were found not only in the genome of B. octagenarum UFV-E1, originally isolated from damaged E. urograndis leaves, but also in the genome of B. octagenarum UFV-OR1, isolated from orchid leaves but not reported as causing damage to its host. However, none of these sequences were located within horizontally transferred regions, suggesting that horizontal gene transfer did not play a significant role in the emergence of the capacity for damaging leaves by Brasilonema. If these proteins also act as virulence factors, leaf damage by this species may be either restricted in host range or triggered by a more complex, uncommon set of factors making it a phenomenon of rare occurrence and thus not commonly observed. Similar sequences were also found in the genomes of B. octagenarum CENA114, obtained from a wet iron water pipe, B. bromeliae SPC 951 and Brasilonema sp. UFV-L1, isolated respectively from bromeliad and ligustrum leaves, which could suggest that the potential for leaf damage is not restricted to a single phyllosphere species, but also present in Brasilonema strains from other habitats and species.

Nevertheless, it is unlikely that these cyanobacteria could be capable of injecting any virulence factors and effectors into their plant hosts considering that the type III secretion system is the main vehicle of infection in known phytopathogenic bacteria but it is absent in this genus. Therefore, even if virulence factors could be expressed by the cyanobacteria in contact with plants, they appear to have no way of reaching the cytoplasm of their hosts since their genomes do not encode the structural proteins necessary for this process. This makes damage by B. octagenarum largely an indirect consequence of the growth of cyanobacterial mats on the leaf surface, causing the blockage of sunlight and hindering photosynthesis by the host, which could possibly progress into more severe consequences after extended colonization. It is therefore possible that other epiphytic cyanobacteria,
even those that do not produce any potential virulence factors, also cause similar damage if specific environmental conditions allow for their biomass to achieve significant volume. Consequently, even though epiphylllic cyanobacteria are traditionally thought of as being universally beneficial to their hosts, benefits to plants may come at a cost that may outbalance them when certain conditions that greatly favor cyanobacterial growth are met.

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Competing Interests

The authors declare there are no competing interests.

Author Contributions

- Danillo O. Alvarenga and Maione W. Franco conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Kaarina Sivonen analyzed the data, authored or reviewed drafts of the paper, and approved the final draft.
- Marli F. Fiore conceived and designed the experiments, analyzed the data, authored or reviewed drafts of the paper, and approved the final draft.
- Alessandro M. Varani conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.

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Data Availability
The following information was supplied regarding data availability:

The sequences are available at GenBank: CP030118–CP030123, QMEA00000000, QMEB00000000, QMEC00000000.

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