Research Article

Influence of Vinasse Application in the Structure and Composition of the Bacterial Community of the Soil under Sugarcane Cultivation

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Although the use of vinasse as a waste helps replenish soil nutrients and improves the quality of the sugarcane crop, it is known that vinasse residues alter the diversity of bacteria naturally present in the soil. The actual impacts of vinasse application on the selection of bacterial taxa are not understood because no studies have addressed this phenomenon directly. Analysis of 16S rRNA gene clone sequences from four soil types showed that the soil planted with sugarcane and fertilized with vinasse has a high diversity of bacteria compared to other biomes, where Acidobacteria were the second most abundant phylum. Although the composition and structure of bacterial communities differ significantly in the four environments (Libshuff’s test), forest soils and soil planted with sugarcane without vinasse fertilizer were similar to each other because they share at least 28 OTUs related to Rhizobiales, which are important agents involved in nitrogen fixation. OTUs belonging to Actinomycetales were detected more often in the soil that had vinasse applied, indicating that these groups are more favored by this type of land management.

1. Introduction

Research on bacterial diversity in soils planted with sugarcane (Saccharum spp.) has indicated that the greater portion of these microorganisms is unknown [1–3]. The importance of the study of bacterial populations in soil is based on approaches depending on the cultivation because these microorganisms could provide new biological resources for the production of commercial inoculants. The bacteria are used in studies that provide insight into mechanisms of the production of compounds that increase sugarcane yield, such as siderophores, indolic components, biological nitrogen fixation, and inorganic phosphate solubilization [3]. Beyond their importance in agronomical applications, Plant Growth-Promoting Bacteria (PGPb) can mitigate the health of contaminated ecosystems when used for bioremediation of a contaminated site [4].

The reuse of agricultural organic waste promotes conservation of natural resources and helps in carbon recycling and recycling of other mineral elements [5, 6]. Based on specific recommendations, reusing these wastes can improve soil quality and reduce dependence on commercial fertilizers that are considered hazardous [7]. Despite its importance for replenishing soil nutrients, vinasse can have a serious environmental impact if used improperly [8] because an excessive accumulation of nutrients can cause contamination [4]. The heterogeneity of vinasse composition contraindicates its use as an organic fertilizer for many reasons, the most common of which is the leaching of potassium nitrate that contaminates the groundwater [7, 9]. Although the effects of fertilization with vinasse on the physicochemical properties of soil have been studied for a long time [7–9], little is known about its impact on the bacterial community.
### 2. Materials and Methods

#### 2.1. Description of the Local Experiments and Soil Samples.

All samples were collected in duplicate at 0–20 cm depth (393 mL). The points were randomly chosen from the central point of each area of interest: (1) soil on which sugarcane was cultivated that was organically fertilized with vinasse (SPV); (2) a heterogeneous soil that contained the master vinasse channel openings were made in the soil and were approximately 15 days because the harvest was finished. The soil was irrigated with vinasse via motorized pumping from the master channel. Due to the constant shifting of vinasse to different cultivation areas when transported in SMC, soil particulates accumulate and form a mixture of different soils with organic waste from the vinasse and crop residues. SFS (475 m in altitude, 2°55′59.90″S and 48°25′49.36″W) was located near the sugarcane cultivation areas, and because it was at a lower altitude, SFS was subject to anthropogenic perturbations of crop management, mainly due to the accumulation of soil and nutrients carried by rainwater. SPS was sampled in January 2013 at the Pau D’alho Farm (Bebedouro City). This area is located at 580 m of altitude (20°55′59.90″S and 48°25′49.36″W) and was planted in sugarcane with manual handling after harvesting the burned sugarcane.

According to the Koeppen climatic classification, Nova Europa and Bebedouro which are located northwest of São Paulo state have warm weather, a tropical rainy climate, a dry winter, and colder months (Aw). Nova Europa city has an annual median temperature of 29.3°C and precipitation of 1,341.4 mm and Bebedouro has a temperature of 31.0°C and precipitation of 1,333.8 mm.

After sampling, a portion of each of the four soils was forwarded to the Laboratory of Analysis of Soils (Faculdade de Ciências Agrárias e Veterinárias de Jaboticabal/SP,unesp, Brazil) to obtain the physical-chemical parameters. The soil properties are shown in Table 1.

#### 2.2. DNA Extraction, PCR, and Cloning.

The total DNA from each soil sample was obtained using a SPIN Kit for Soil* (MP Biomedicals), according to the manufacturer’s instructions. 16S rRNA gene was partially amplified (variable regions 1 and 2) with Y1 and Y2 primers [16], yielding Polymerase Chain Reaction (PCR) products of approximately 300 bp. The PCR reaction mixture consisted of 20–50 ng/µL of DNA, 5.0 pmol/µL of each primer, 0.2 mM of dNTPs, 3.0 mM of MgCl₂, Buffer 1x, and 2.5 U Taq DNA polymerase (Ludwig Biotec). A thermocycler model PTC-100™ Programmable Thermal Controller (MJ Research, Inc.) used a thermal profile of 95°C for 2 min, 35 cycles of 95°C for 45 s, 65°C for 45 s, and 72°C for 1 min, and 30 s ending with 72°C for 5 min. After the PCR reaction, the products were purified with a Wizard* SV Gel and PCR Clean-Up System (Promega) and for PCR

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#### Table 1: Physicochemical properties of the soil from four biomes studied in this work at depth of 0–20 cm.

| Soils  | pH | OM g/dm³ | P resin mg/dm³ | K mmol/dm³ | Ca mmol/dm³ | Mg mmol/dm³ | H + AL mmol/dm³ | SB mmol/dm³ | T % |
|-------|----|----------|----------------|------------|-------------|-------------|----------------|-------------|-----|
| SPV   | 5.5| 20       | 33             | 9.8        | 44          | 19          | 31             | 72.8        | 103.8| 70 |
| SMC   | 6.4| 19       | 26             | 10.3       | 106         | 26          | 12             | 142.3       | 154.3| 92 |
| SPS   | 4.8| 10       | 9              | 1.1        | 39          | 22          | 38             | 62.1        | 100.1| 62 |
| SFS   | 5.2| 23       | 31             | 1.1        | 39          | 22          | 38             | 62.1        | 100.1| 62 |

pH: hydrogen potential; OM: organic matter; K: potassium; Ca: calcium; Mg: magnesium; H + AL: exchangeable acidity or potential acid; SB: sum of bases; T: Cation Exchange Capacity (CEC) effective at pH 7.0; V: percentage of base saturation; mmol/dm³: millimol per cubic decimeter.

SPV: soil with stillage fertigation and sugarcane cultivation; SMC: heterogeneous soil deposited on the stillage master channel; SPS: soil by sugarcane planting without stillage fertigation; SFS: native forest soil located near areas of sugarcane cultivation.

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Such research has been nonspecific because indirect techniques to assess local microbiota such as respirometry are generally used [10]. It is commonly known that the use of vinasse, compared to the application of other agricultural wastes, contributes to improvement of the biological quality of the soil because of an increase in the total bacterial population, in particular of group Actinomycetes [7].

The use of the molecular techniques based on 16s ribosomal RNA (16s rRNA) gene clone library allows an indirect assessment of the bacterial community, composing the bacterial community of soils subjected to different agricultural management practices, and of the impacts on the composition and diversity of different bacterial groups [1, 2, 11]. Considering that the major part of the bacterial population in soil cannot be cultivated by traditional techniques [5, 12] and play a crucial role in the maintenance of different ecosystems [2, 13–15], these microorganisms are considered as biological indicators of soil quality [7].

This study aimed to identify the bacterial communities in soils in which sugarcane was cultivated and subject to different management practices using the analysis of partial sequences of clones from 16s rRNA gene libraries.
cloning CloneJET™ Cloning Kit (Fermentas) was used as per
the manufacturer's instructions.

Colonies in Petri dishes were collected with a sterile wooden toothpick and organized into 96-well ELISA plates containing 100 µL of Luria-Bertani medium with ampicillin (50 µg/mL). After overnight development of the clones at 37°C, a copy of each library was inoculated to extract the plasmid DNA [17] and 100 µL of 40% glycerol (v/v) was added to the original library for storage at −80°C.

2.3. Sequencing and Taxonomic Evaluation. Amplicons of 16S rRNA gene were sequenced using Y1 primer. DNA sequences were obtained with ABI PRISM Big Dye Terminator cycle sequencing-ready reaction kit (Applied Biosystems) using ABI 3100 (Perkin Elmer) capillary sequencer, following the manufacturer's instructions. The sequences used for the analyses were those with quality Phred ≥ 20 [18]. A search for chimeric sequences was performed with Decipher software [19] and taxonomic identification to the level of phylum, class, order, family, and genus was made with the Classifier software using confidence limits of 80, 85, 91, 92, and 95% [11], respectively. Sequences that scored 95% confidence in the Classifier were used to search for similar sequences in MegaBlast (16S rRNA gene of the Bacterial and Archaea) [20] from the National Center for Biotechnology Information (NCBI) or the SeqMatch Ribosomal Database Project (RDP II) [21]. These searches were conducted in March 2016.

2.4. Construction of the Classification Phylogenetic Tree. The sequences of 16S rRNA gene were submitted to GenBank under the following accession numbers: SPV (KJ749164–KJ749443), SMC (KJ749444–KJ749650), SPS (KJ748893–KJ749163), and SFS (KJ748700–KJ748892). The sequences from the four ecosystems (SPV, SMC, SPS, and SFS) (MegaBlast or SeqMatch) were used to construct the phylogenetic tree. The database sequences that were most similar to the sequences of clones from this study were used to compare the phylogenetic tree after recovery from the GenBank/National Center for Biotechnology Information (NCBI). Alignment was performed based on ClustalW method [22] available on BioEdit 7.2.3 [23]. The distance matrices were calculated from the sequence alignment program using Dnadist from the package program PHYLIP v3.6 [24] using the matrix of nucleotide substitutions of Jukes and Cantor [25] for the correction of mutations. MEGA 6.06 [26], a neighbor-joining method [27], bootstrapping with 1,000 replications and an optional pairwise deletion, was used for the construction of the phylogenetic tree.

2.5. Diversity Estimation, Comparison of the Library, and Statistical Analysis. α-diversity was obtained by analyzing the sequences of all clones (SPV, SMC, SPS, and SFS) followed by the construction of a rarefaction curve and obtaining the indices of richness, evenness, Shannon (H), and Chao 1, considering an evolutionary distance of 3% difference (or 97% similarity). H was also used to estimate γ-diversity among the samples. Significant differences among the structure of the bacterial communities were indicated by the Libshuff test (β-diversity) using the Cramer-von Mises criterion (P ≤ 0.05). P value obtained was corrected with the Bonferroni correction. All data were obtained with mothur v.1.33.3 [28]. UniFrac [29] analysis was used to compare the clones of each library using a normalization step with 100 permutations, the Jackknife Environment Clusters option, and the considerate selection algorithm, which considered the relative abundance of the OTUs.

3. Results

3.1. Phylogenetic Groups Identified in Various Environments. A total of 951 partial sequences of 16S rRNA genes (free of chimeric sequences) were obtained, of which 280 were recovered from the soil on which sugarcane was cultivated and fertilized with vinasse (SPV), 271 sequences from soil on which sugarcane was cultivated (SPS), 207 from the master vinasse channel (SMC), and 193 sequences from the secondary forest (SFS). All sequences of 16S rRNA gene amplicons were combined and used for subsequent analyses.

The taxonomic classification in RDP II indicated 11 different phylogenetic groups: SPV had II phyla, SPS had seven phyla, SMC had five phyla, and SFS had four phyla (Table 2). Proteobacteria were the most abundant phylum in the four soil types and were more frequent in SPS (63%) and SFS (60%), where Alphaproteobacteria occurred at more than 50% frequency. A few relative sequences (less than 5%) of Betaproteobacteria were identified. Gammaproteobacteria were detected only in SPS and SMC, and Deltaproteobacteria were only in SPV and SFS.

Actinobacteria and Acidobacteria were the second and third most abundant phyla in SPS and SFS, as reported in other studies [30]. Different than observed in other soils, the second most abundant phylum in SPV was Acidobacteria (28%), followed by Actinobacteria at 6% of the taxonomic affiliations. Actinobacteria and Acidobacteria showed same abundance (13-14%) in SMC; and Firmicutes (7%) phylum had the third highest frequency. Although not shared by all environments, less frequent phyla were also identified in the four environments, such as Planctomycetes, Nitrospirae, Bacteroidetes, Chloroflexi, Armatimonadetes, and TM7 ("Saccharibacteria") (Table 2). Some clones (17% total) showed no similarity more than 80% with available sequences in RDP II; for this reason, this group was assigned to the unknown bacteria. The SPS ecosystem had fewer affiliated sequences as unclassified bacteria (5%), while the SMC was the environment showing the high number of OTUs for that group (25%) (Table 2).

A high frequency of OTUs related to order Rhizobiales (Alphaproteobacteria) was found, which is a group of bacteria commonly found in soil that play an important role in biological nitrogen fixation [11]. Order Rhizobiales had different frequencies across the four biomes, identified in 109 OTUs at SPS, 73 OTUs at SFS, 39 OTUs at SPV, and 19 OTUs at SMC. All environments showed a higher taxon (Classifier), although most similar sequences were uncultured bacteria obtained from the analysis of 16S rRNA genes in the soil recovered (Figures 1 and 2).
Table 2: Bacterial phylogenotypes distribution in clone libraries of 16S rRNA gene as affiliation by Classifier algorithm available on RDP II using cutting 80% confidence.

| Phylogenetic groups                  | SPV | SPS | SMC | SFS | Total |
|--------------------------------------|-----|-----|-----|-----|-------|
| Proteobacteria                       | 30.3| 62.9| 40.0| 59.6| 48.2  |
| Alphaproteobacteria                  | 22.1| 56.8| 19.3| 55.0| 38.3  |
| Betaproteobacteria                   | 5.0 | 1.8 | 0.4 | 0.4 | 2.0   |
| Gammaproteobacteria                  | —   | 0.7 | 6.4 | —   | 1.8   |
| Deltaproteobacteria                  | 0.4 | —   | —   | 0.4 | 0.2   |
| Unclassified Proteobacteria          | 2.9 | 3.2 | 13.9| 3.6 | 6.0   |
| Acidobacteria                        | 28.2| 8.9 | 13.2| 6.1 | 14.1  |
| Actinobacteria                       | 6.4 | 20.7| 13.9| 14.6| 13.9  |
| Firmicutes                           | 2.9 | 0.7 | 7.1 | —   | 2.7   |
| Gemmatimonadetes                     | 1.4 | 0.7 | 0.3 | 2.5 | 1.3   |
| Othersb                              | 11.0| 1.0 | —   | —   | 2.9   |
| Unclassified bacteria                | 20.0| 5.0 | 2.5 | 17.1| 17    |

a Abbreviations of the four biomes are shown in Table 1.
b Others: Planctomycetes, Nitrospirae, Bacteroidetes, Chloroflexi, Armatimonadetes, Candidatus phyla Saccharibacteria, and Candidatus phyla division WPS-1.

Normalization factor is 280/total, where total is sum of the number of sequences not chimeras. The normalization factor of each environment was multiplied by each taxon observed at the level of phylum and converted to percentage.

For all soils, the formation of clusters related to subdivisions 1–6, 11, 13, and 16 Acidobacteria species in RDP II (Classifier and SeqMatch) was observed. Similar to order Rhizobiales, Gp3 and Gp4 subgroups of Acidobacteria appeared to be more sensitive to the relative frequency; as SPV represented 22–32 OTUs affiliated to species of this group, SPS, SFS, and SMC had 2–4 species (Figure 1).

3.2. Diversity of Bacterial Communities in Environments. The diversity environmental indices were obtained using OTUs with an evolutionary distance ≤ 0.03. All of the diversity and richness indices (Table 4), supported by the slope of the rarefaction curve (data not shown), showed the following tendency in terms of species: SPV > SPS > SFS > SMC. Although the rarefaction curve for SFS and SPS seemed to indicate a similar profile, SPS was considered to be an environment with intermediate diversity according to values obtained by Shannon and Simpson and richness estimator Chao 1 (less abundant acceptable OTUs). The less diverse environment was the SMC biome according to the diversity indices (Table 4). Because the rarefaction curve did not plateau and the slope was high for all environments (data not shown), it is evident that many new taxa could still be identified in any of these ecosystems, corroborated by the number of OTUs observed (S) and the values obtained by estimator Chao 1 (Table 4).

3.3. Effects of the Abundance of OTUs in the Structure of Bacterial Communities. The UniFrac Jackknife Environmental Cluster analysis clearly shows that SPS and SFS share few taxa in common comparing the four environments (Figure 3). For 10 shared OTUs identified by mothur (97% similarity) in those environments, the relationship was almost exclusive to Alphaproteobacteria, especially those of order Rhizobiales (Tables 3 and 5). In the latter test, SMC and SPV appear to contain bacterial communities distinct from those found in SFS and SPS. Although SFS and SPS seemed to be relatively similar (Figure 3), the Libshuff molecular variance test (data not shown) indicated significant differences (P < 0.01). In this test, the four bacterial communities recovered from soils in directions x-y and y-x demonstrated that the sequences from each library are quite different in each type of environment.

4. Discussion

4.1. Distinction of Phylogenetic Groups Identified in Soils. Proteobacteria, a group with great morphological, physiological, and metabolic variation [31], have been found as the most abundant group in soil [30] and play an important role in the global carbon, nitrogen, iron, and sulfur cycles [5, 6, 32]. This bacterial group has been frequently reported in soils planted with sugarcane using cultivation-dependent techniques [3] and also by cultivation-independent molecular techniques [1, 2]. Although many proteobacterial strains have been isolated and characterized, an analysis based on 16S rRNA gene sequences from environmental clones and by pyrosequencing indicated that many members are still unknown, and, therefore, their specific roles in various ecosystems (marine sediments, soil, anaerobic degradation of organic components, etc.) are also unknown [5].

In our research, we found that the frequency of Proteobacteria and Actinobacteria phyla and their classes was variable depending on the type of soil management. SPV was the environment that showed the lowest frequency of those Proteobacteria in relation to SPS and SFS (Table 2). Those data demonstrate that Proteobacteria presented impaired abundance in soil under vinasse application (SPV) compared with that under burning management (SPS), in spite of the application of vinasse as a source of essential nutrients.
Figure 1: Phylogenetic tree showing species of the Acidobacteria group identified four biomes. The access numbers to the GenBank sequences obtained in this study are shown in parentheses. The numbers in brackets represent the number of sequences that were affiliated to the same taxonomic group Acidobacteria. For the tree construction was used the matrix of Jukes and Cantor nucleotide substitutions, neighbor-joining method with bootstrap for 1,000 replicas and pairwise deletion option. The nodes showed just bootstrap value of above 50%. The scale indicates 0.05 occurring nucleotide substitutions at each position. The abbreviations of the four biomes are shown in Table 1.
to sugarcane crop. With the exception of SMC, all of the classes of Proteobacteria had sequences that conformed to those reported in most studies on bacterial diversity in soil [11]. These observations are important because we need to document as much information as possible about the effects of vinasse on soil microbiota, once those evidences support the development of new researches and the improvement of more sustainable agricultural techniques than those available. Thus, one can promote the increase of crop productivity and improve the use of byproducts generated by sugarcane industry.

We also observed, in the four environments, sequences associated with clones assigned to Alpha-, Beta-, Gamma-, and Deltaproteobacteria (Table 2), which emphasizes that
Each of the ecosystems has different sources for the replacement of organic matter such as natural vegetation debris (litter in SFS), crop residues from other crops (SPV, SFS, and SMC), and vinasse (SPV). Probably because of this wide variation in organic matter inputs, the Actinobacteria phylum was detected as one of the three most abundant phyla in these environments, appearing to be abundant in SPV. We also emphasize that almost all of the OTUs related to Actinobacteria from the four biomes were similar to bacterial strains commonly isolated from soil, showing many well-known characterized strains (Figure 2).

In soil, the population growth of heterotrophic bacteria is attributed to the presence of vinasse [10]. Our results agree with these predictions because many heterotrophic bacterial groups were identified within 11 phyla identified in SPV, such as Proteobacteria, Acidobacteria, Actinobacteria, Firmicutes, and Bacteroidetes (Table 2). These microorganisms have been described as promoters of soil quality control, where the vinasse has potential to select bacterial taxa such as Actinomycetes [7]. In SPV, Actinobacteria (or Actinomycetes) had a higher frequency of OTUs related to the Micromonosporaceae, Nocardioidaceae, Mycobacteriaceae, and Pseudonocardiaee families (partially shown in Figure 2), that seem to be more favored in SPV than in the other three biomes.

4.2. Richness and Diversity of the Four Biomes. In relation to previous studies in subtropical forest soils [14], our ecosystems were classified as having high diversity. According to analyses of indirectly accessed soil microorganisms in the presence of vinasse [7, 10], SPV had a higher bacterial diversity than any other biome analyzed in this study, showing the potential of vinasse to nurture plenty of bacterial species in the soil (Table 4 and Figure 1). Forest soils have less diversity of bacterial taxa than agricultural soils, a fact related to the increase or absence of the abundance of certain bacterial taxa in forest soils for agricultural soils [11]. For these reasons, SFS showed less diversity in bacterial species than the other three soils studied, with the exception of SMC (Table 4), probably due to the very heterogeneous conditions to which that environment was subjected.

4.3. Differences in the Structure of Bacterial Communities. Changes caused by different types of natural and/or anthropogenic disturbances on agricultural land and in forests cause significant changes in the composition of the bacterial communities [11] including the effects of sugarcane transgenic cultivation and herbicide application on bacterial communities in soil [1]. The Libshuff test demonstrated these perturbations, and significance of $P < 0.01$ (data not shown) was obtained between the four environments. Even with differences in the composition of the bacterial communities, SPV and SFS shared some bacterial taxa (Table 3), indicating that although subject to different management practices, some taxa that compose the bacterial communities in these soils showed physiological and metabolic similar characteristics (Figure 3), as OTUs belonging to order Rhizobiales.

Those differences between the bacterial communities for the four environments directly reflected the observed
Table 3: OTUs of Alphaproteobacteria formed with mothur (3% similarities) and identified by MegaBlast and Classifier from RDP II.

| OTU formed | Shared by clones environments | Description based on RDP II classification* | Accession (16S rRNA database) | Results in MegaBlast Similarities in MegaBlast (%) | Genus |
|------------|--------------------------------|-----------------------------------------------|-----------------------------|-----------------------------------------------|-------|
| Otu001     | SFS004, SFS058, SFS189, SPS048, and SPS052 | Order Rhizobiales and family Hyphomicrobiaceae unclassified (SFS004, SFS058, SFS189, and SPS052) and *Rhodoplanes* (SPS048) | NR_134225.1 | 97 | *Varibacter gotjawalensis* |
| Otu004     | SPS074, SPS064, SPS075, SFS037, and SPV011 | Order Rhizobiales (SPS075 and SFS037) and family Hyphomicrobiaceae unclassified (SPS074, SPS064, and SPV011) | NR_125638.1 | 96 | *Bradyrhizobium valentinum* |
| Otu011     | SFS036, SFS027, and SPS171 | Order Rhizobiales (genus *Nitrobacter* SFS036 and SFS027) and Bradyrhizobiaceae unclassified (SPS171) | NR_125638.1 | 100 | *Bradyrhizobium valentinum* |
| Otu012     | SFSI22, SFS193, and SPS012 | Order Rhizobiales and Xanthobacteraceae unclassified (SFSI22) and Rhizobiales unclassified (SFS193 and SPS012) | NR_043515.1 | 95 | *Pseudolabrys taiwanensis* |
| Otu018     | SFS057 and SPS023 | Family Bradyrhizobiaceae (SPS023) and genus *Bradyrhizobium* (SFS057) | NR_133987.1 | 99 | *Bradyrhizobium neotropicalis* |
| Otu019     | SFS069 and SPS024 | Order Sphingomonadaceae (SPS069) and family Sphingomonadaceae (SPS024) | NR_133862.1 | 97 | *Sphingomonas daechungensis* |
| Otu039     | SFS064 and SPS173 | Order Rhizobiales (SPS173) and Methyllocystaceae unclassified (SFS064) | NR_134515.6.2 | 94 | *Rhodoplanes oryzae* |
| Otu052     | SFS025 and SPS203 | Order Rhizobiales (SPS025) and Hyphomicrobiaceae unclassified (SPS203) | NR_134225.1 | 96 | *Varibacter gotjawalensis* |
| Otu067     | SFS071 and SPS080 | Family Sphingomonadaceae (SPS080) and Sphingomonadaceae (SPS071) | NR_074280.1 | 96 | *Sphingopyxis alaskensis* |
| Otu068     | SFS182 and SPS078 | Order Rhizobiales (SPS078) and Rhizobiales unclassified (SFS182) | NR_112190.1 | 94 | "*Rhodoplanes cryptolactis*" |

*Based on 95% confidence threshold in Classifier.
Table 4: Diversity and richness rates observed for each bacterial community present in the soils analyzed. The results were based on 16S rRNA gene of the bacterial clones obtained from 4 environments.

| Indices | SPV | SMC | SPS | SFS |
|---------|-----|-----|-----|-----|
| S       | 188 | 168 | 176 | 179 |
| N       | 193 | 193 | 193 | 193 |
| Evenness| 0.98| 0.97| 0.97| 0.97|
| Richness| 0.93| 0.76| 0.83| 0.85|
| Shannon  | 5.23 (5.33) | 5.06 (5.17) | 5.13 (5.24) | 5.14 (5.25) |
| Simpson  | 0.0003 | 0.0019 | 0.0001 | 0.0012 |
| Chao 1   | 2,963 (1,488–6,109) | 1,039 (645–1,758) | 1,262 (765–2,178) | 2,231 (1,172–4,415) |

a Calculations were based on OTUs formation based on evolutionary distance of ≤ 0.03. b Abbreviations are shown in Table 1. c S is number of OTUs observed. d N is sequences number. e Evenness is $H/H_{\text{max}}$ [44]. f Richness is (number of OTUs singletons + 1)/log(N). The observed value/maximum possible is informed. g Maximum value observed for Shannon ($H$). h Confidence intervals (95%) for Chao 1. i Maximum value observed for singletons.

Table 5: OTUs related to bacterial orders in four soil biomes (cutoff of 95% confidence in RDP II Classifier) and identity by MegaBlast.

| Sequence | Genus | Accession GenBank | Similarities (%) | Description | References |
|----------|-------|-------------------|------------------|-------------|------------|
| SPV212   | *Azohydromonas lata* | NR_041244.1 | 94 | Nitrogen fixation in different soils | [38, 45] |
| SPV150   | *Cupriavidus metallidurans* | NR_027607.1 | 97 | Chemolithotrophic bacteria common in agricultural soils and showing potential for bioremediation of soils contaminated with Cd, Zn, and Cu, among others | [39, 46] |
| SPV115   | *Bradyrhizobium valentinum* | NR_125638.1 | 97 | Nitrogen fixation in different soils | [40] |
| SPS230   | *Caulobacter vibrioides* | NR_037099.1 | 97 | Utilizes various sources of carbon and has potential for application in selenium solubilizing | [42, 47] |
| SPS052   | “Rhodoplanes cryptolactis” | NR_112190.1 | 96 | Nitrogen fixation, organic matter decomposition, and growth promotion in plants | [43] |
| SMC121   | *Acidiphilium cryptum* | NR_119294.1 | 91 | Iron reduction (Fe$^{3+}$ to Fe$^{2+}$) in acidic conditions | [41] |
| SFS043   | *Sphingosinicella microcystinivorans* | NR_040927.1 | 96 | Relatively new genus which has the capacity to degrade polyaromatic hydrocarbons | [48] |

bacterial species. In particular, the species of Proteobacteria were the most diverse and environmental specific, in which the clones showed similarity with bacterial isolates from different important environmental physiological profiles (Table 5). In SPV, *Azohydromonas lata* (SPV212), *Cupriavidus metallidurans* (SPV150), and *Bradyrhizobium valentinum* (SPV115) are generally associated with nitrogen fixation and with great potential for use in bioremediation of sites contaminated with heavy metals [38–40]. *Acidiphilium cryptum* (SMC121) identified in SMC is associated with more acidic environments and possesses the ability to reduce Fe$^{3+}$ and Fe$^{2+}$, denoting participation in the iron cycle in soils [41]. SPS presented *Caulobacter vibrioides* (SPS230) and *Rhodoplanes cryptolactis* (SPS052) participating in the carbon sources metabolism arising from organic matter in the soil, besides nitrogen fixation and growth promotion in plants [42, 43]. This comparison shows that, in nonadded vinasse soil (SPS), species with various metabolic profiles and more adapted to plant-interaction seems to be favored. However, the SPV and SMC soils present greater availability of macro- and micronutrients than other soils probably due to the wealth of organic matter and minerals present in vinasse [7–10], with exceptions of nitrogen and iron. For this reason, to address the lack of these nutrients, we speculate that bacteria are playing major important ecological roles in nitrogen and iron cycles and they are even favored in the presence of vinasse.

5. Conclusion

The vinasse increases bacterial diversity in soil and promotes favoring species participating in the nitrogen and iron cycle. Species subdivisions 3 and 4 of Acidobacteria were more abundant in soil in which vinasse was applied. Species of
families belonging to Actinomycetales were more diverse in soil in which vinasse had been applied for sugarcane cultivation than in other soils.

Competing Interests
The authors declare no competing interests regarding the publication of this paper.

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References
[1] F. Dini-Andreote, F. D. Andreote, R. Costa, R. G. Taketani, J. D. van Elsas, and W. L. Araújo, "Bacterial soil community in a Brazilian sugarcane field," Plant and Soil, vol. 336, no. 1-2, pp. 337–349, 2010.
[2] G. Pisa, G. S. Magnani, H. Weber et al., "Diversity of 16S rRNA genes from bacteria of sugarcane rhizosphere soil," Brazilian Journal of Medical and Biological Research, vol. 44, no. 12, pp. 1215–1221, 2011.
[3] A. Beneduzi, F. Moreira, P. B. Costa et al., "Diversity and plant growth promoting evaluation abilities of bacteria isolated from sugarcane cultivated in the South of Brazil," Applied Soil Ecology, vol. 63, pp. 94–104, 2013.
[4] L. E. de-Bashan, J.-P. Hernandez, and Y. Bashan, "The potential contribution of plant growth-promoting bacteria to reduce environmental degradation—a comprehensive evaluation," Applied Soil Ecology, vol. 61, pp. 171–189, 2012.
[5] A. M. Spain, L. R. Krumholz, and M. S. Elshahed, "Abundance, composition, diversity and novelty of soil Proteobacteria," The ISME Journal, vol. 3, no. 8, pp. 992–1000, 2009.
[6] S. Hedrich, M. Schlömann, and D. B. Johnson, "The iron-oxidizing proteobacteria," Microbiology, vol. 157, no. 6, pp. 1551–1564, 2011.
[7] R. D. M. Prado, G. Caione, and C. N. S. Campos, "Filter cake and vinasse as fertilizers contributing to conservation agriculture," Applied and Environmental Soil Science, vol. 2013, Article ID 581984, 8 pages, 2013.
[8] V. Parneau, N. Condom, R. Oliver, P. Cazeville, and S. Recous, "Vinasse organic matter quality and mineralization potential, as influenced by raw material, fermentation and concentration processes," Bioresource Technology, vol. 99, no. 6, pp. 1553–1562, 2008.
[9] M. B. Benke, A. R. Mermut, and H. Shariatmadari, "Retention of dissolved organic carbon from vinasse by a tropical soil, kaolinite, and Fe-oxides," Geoderma, vol. 91, no. 1-2, pp. 47–63, 1999.
[10] A. P. Mariano, S. H. R. Crivelaro, D. D. F. de Angelis, and D. M. Bonotto, "The use of vinasse as an amendment to ex situ bioremediation of soil and groundwater contaminated with diesel oil," Brazilian Archives of Biology and Technology, vol. 52, no. 4, pp. 1043–1055, 2009.
[11] R. Upchurch, C.-Y. Chiu, K. Everett, G. Dyzsynski, D. C. Coleman, and W. B. Whitman, "Differences in the composition and diversity of bacterial communities from agricultural and forest soils," Soil Biology and Biochemistry, vol. 40, no. 6, pp. 1294–1305, 2008.
[12] P. D. Schloss and J. Handelsman, “Toward a census of bacteria in soil,” PLoS Computational Biology, vol. 2, no. 7, article e92, 2006.
[13] O. Nercessian, Y. Fouquet, C. Pierre, D. Prieur, and C. Jeannin, "Diversity of Bacteria and Archaea associated with a carbonate-rich metalliferous sediment sample from the Rainbow vent field on the Mid-Atlantic Ridge," Environmental Microbiology, vol. 7, no. 5, pp. 698–714, 2005.
[14] Y.-T. Lin, W. B. Whitman, D. C. Coleman, and C.-Y. Chiu, "Comparison of soil bacterial communities between coastal and inland forests in a subtropical area," Applied Soil Ecology, vol. 60, pp. 49–55, 2012.
[15] L. F. Martins, L. P. Antunes, R. C. Pascon et al., "Metagenomic analysis of a tropical composting operation at the São Paulo Zoo Park reveals diversity of biomass degradation functions and organisms," PLoS ONE, vol. 8, no. 4, Article ID e61928, 2013.
[16] J. P. W. Young, H. L. Downer, and B. D. Eardly, "Phylogeny of the phototrophic rhizobium strain BTAi1 by polymerase chain reaction-based sequencing of a 16S rRNA gene segment," Journal of Bacteriology, vol. 173, no. 7, pp. 2271–2277, 1991.
[17] J. Sambrook and D. W. Russell, Molecular Cloning: A Laboratory Manual, vol. 1 of Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, NY, USA, 3rd edition, 2001.
[18] D. Gordon, C. Abajian, and P. Green, "Consed: a graphical tool for sequence finishing," Genome Research, vol. 8, no. 3, pp. 195–202, 1998.
[19] E. S. Wright, L. S. Yilmaz, and D. R. Noguera, "DECIPHER, a search-based approach to chimera identification for 16S rRNA sequences," Applied and Environmental Microbiology, vol. 78, no. 3, pp. 717–725, 2012.
[20] A. Morgulis, G. Coulouris, Y. Raytselis, T. L. Madden, R. Argarwala, and A. A. Schäffer, "Database indexing for production MegaBLAST searches," Bioinformatics, vol. 24, no. 16, pp. 1757–1764, 2008.
[21] J. R. Cole, Q. Wang, J. A. Fish et al., "Ribosomal database project: data and tools for high throughput 16S RNA analysis," Nucleic Acids Research, vol. 42, no. 1, pp. D633–D642, 2014.
[22] J. D. Thompson, D. G. Higgins, and T. J. Gibson, "CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice," Nucleic Acids Research, vol. 22, no. 22, pp. 4673–4680, 1994.
[23] T. A. Hall, "BioEdit: a user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT," Nucleic Acids Research, vol. 41, pp. 95–98, 1999.
[24] J. Felsenstein, PHYLIP (Phylogeny Inference Package) Version 3.6, Department of Genome Sciences and Department of Biology, University of Washington, Seattle, Wash, USA, 2005.
[25] T. H. Jukes and C. R. Cantor, "Evolution of protein molecules," in Mammalian Protein Metabolism, H. N. Munro, Ed., pp. 21–132, Academic Press, New York, NY, USA, 1969.
[26] K. Tamura, G. Stecher, D. Peterson, A. Filipski, and S. Kumar, "MEGA6: molecular evolutionary genetics analysis version 6.0," Molecular Biology and Evolution, vol. 30, no. 12, pp. 2725–2729, 2013.
[27] N. Saitou and M. Nei, "The neighbor-joining method: a new method for reconstructing phylogenetic trees," Molecular Biology and Evolution, vol. 4, no. 4, pp. 406–425, 1987.
28] P. D. Schloss, S. L. Westcott, T. Ryabin et al., “Introducing mothur: open-source, platform-independent, community-supported software for describing and comparing microbial communities,” Applied and Environmental Microbiology, vol. 75, no. 23, pp. 7535–7541, 2009.
29] C. Lozupone, M. Hamady, and R. Knight, “UniFrac—an online tool for comparing microbial community diversity in a phylogenetic context,” BMC Bioinformatics, vol. 7, no. 1, article 371, pp. 1–14, 2006.
30] P. H. Janssen, “Identifying the dominant soil bacterial taxa in libraries of 16S rRNA and 16S rRNA genes,” Applied and Environmental Microbiology, vol. 72, no. 3, pp. 1719–1728, 2006.
31] M. Ros, B. A. Knapp, U. Peintner, and H. Insam, “Short communication. Response of bacterial community composition to long-term applications of different composts in agricultural soils,” Spanish Journal of Agricultural Research, vol. 9, no. 1, pp. 338–343, 2011.
32] R. F. Fulthorpe, L. F. W. Roesch, A. Riva, and E. W. Triplett, “Distantly sampled soils carry few species in common,” The ISME Journal, vol. 2, no. 9, pp. 901–910, 2008.
33] M. Yoshida, S. Ishii, D. Fujii, S. Otsuka, and K. Senoo, “Identification of active denitrifiers in rice paddy soil by DNA- and RNA-based analyses,” Microbes and Environments, vol. 27, no. 4, pp. 456–461, 2012.
34] X. Chen, Y. Su, X. He, Y. Wei, W. Wei, and J. Wu, “Soil bacterial community composition and diversity respond to cultivation in Karst ecosystems,” World Journal of Microbiology and Biotechnology, vol. 28, no. 1, pp. 205–213, 2012.
35] M. K. Männistö, M. Tiirola, and M. M. Häggblom, “Bacterial communities in Arctic fjuals of Finnish lapland are stable but highly pH-dependent,” FEMS Microbiology Ecology, vol. 59, no. 2, pp. 452–465, 2007.
36] Y.-T. Lin, Y.-J. Huang, S.-L. Tang, W. B. Whitman, D. C. Coleman, and C.-Y. Chiu, “Bacterial community diversity in undisturbed perhumid montane forest soils in Taiwan,” Microbial Ecology, vol. 59, no. 2, pp. 369–378, 2010.
37] J. F. Araújo, A. P. de Castro, M. M. C. Costa et al., “Characterization of soil bacterial assemblies in Brazilian savanna-like vegetation reveals Acidobacteria dominance,” Microbial Ecology, vol. 64, no. 3, pp. 760–770, 2012.
38] M. C. P. Silva, B. Schloter-Hai, M. Schloter, J. D. van Elsas, and J. F. Salles, “Temporal dynamics of abundance and composition of nitrogen-fixing communities across agricultural soils,” PLoS ONE, vol. 8, no. 9, Article ID e74500, 2013.
39] L. Diels, S. Van Roy, S. Taghavi, and R. Van Houdt, “From industrial sites to environmental applications with Cupriavidus metallidurans,” Antonie Van Leeuwenhoek, vol. 96, no. 2, pp. 247–258, 2009.
40] D. Durán, L. Rey, A. Navarro, A. Busquets, J. Imperial, and T. Ruiz-Argüeso, “Bradyrhizobium valentinum sp. nov., isolated from effective nodules of Lupinus mariae-josephae, a lupine endemic of basic-lime soils in Eastern Spain,” Systematic and Applied Microbiology, vol. 37, no. 5, pp. 336–341, 2014.
41] K. Küsel, T. Dorsch, G. Acker, and E. Stackebrandt, “Microbial reduction of Fe(III) in acidic sediments: isolation of Acidiphilium cryptum JF-5 capable of coupling the reduction of Fe(III) to the oxidation of glucose,” Applied and Environmental Microbiology, vol. 65, no. 8, pp. 3633–3640, 1999.
42] I. I. Berestovskaia, A. M. Lysenko, T. P. Turova, and L. V. Vasili’eva, “A psychrotolerant Caulobacter sp. from Russian polar tundra soil,” Mikrobiologija, vol. 75, no. 3, pp. 377–382, 2006.