INTRODUCTION

Agricultural soils have a microbial diversity that can degrade if under stressful conditions for a prolonged period of time. Therefore, the isolation of microorganisms is beneficial for their use as inoculants to increase the degradation of contaminants (VILLAVERDE et al., 2018).

Soil bio-augmentation, strain introduction, and microbial consortium with the desired catalytic abilities increases the degradation capacity of xenobiotics (MROZIK et al., 2010). Therefore, phytoremediation associated with microorganisms comprises a promising technology for environmental decontamination (SALT et al., 1998). Bacteria can interact with the contaminated environment in different ways, either by promoting plant growth, which favors plants under stress, degrading toxic compounds, or increasing nutrient availability.

Hexazinone is an herbicide of the triazinone chemical group used in sugarcane crops (TONIETO & REGINATO, 2014). According to
MARTINS (2015), the study and assessment of the risks of this compound in the environment are relevant because it presents moderate solubility in an aqueous medium. Contamination can occur in a variety of media, including water, soil, and air, resulting in potential risks. The main degradation route of this herbicide in soil is attributed to microorganisms. However, the degradation rate is slow, and the residual activity can extend over several months. WANG et al. (2012) state that the degradation of this molecule is associated with organic carbon content and microbial activity in the soil.

This study selected hexazinone-tolerant microorganisms with phosphate solubilization capacity as well as prepared inoculants to be used in association with plants in the remediation of soil contaminated by hexazinone.

MATERIALS AND METHODS

Isolation, DNA extraction, and identification

First, 20 soil samples were collected at a depth of 10 cm from Rio Verde, GO, at a location (17º 48' 28.2" South and 50º 54' 39" West, altitude 720 m) with a history of hexazinone application. Samples were homogenized to obtain a composite sample. The microorganisms were isolated from serial dilution, plated from 1 g of soil, and diluted in 9 mL of saline solution (0.85%).

Plating was performed with 0.1 mL of each sample being spread on Petri dishes containing nutrient medium (rich medium) with and without herbicide and minimum medium [(NH₄)₂SO₄, 1 g; NaCl, 1 g; KH₂SO₄, 1.5 g; MgSO₄, 0.2 g; agar, 15 g] with and without herbicide. The treatments without herbicide served as controls. Treatments were entirely randomized in a 4 × 9 factorial scheme, consisting of the rich medium with and without herbicide, the minimum medium with and without herbicide, and nine dilutions (up to 10⁹) with three repetitions.

The herbicide concentration was 4.86 mL L⁻¹. Plates were incubated at 28 ºC for 24 and 48 h for rich and minimum medium, respectively, following which the colony forming units (CFUs) were counted. Hexazinone-tolerant bacterial colonies were identified by the presence of a degradation halo and based on the morphological differences observed in the plates containing the herbicide.

The 16S rDNA gene was amplified by polymerase chain reaction (PCR) using R1387 (CGG TGT GTA CAA GGC GGA ACG) and POF27F (GAG AGT TTG ATC CTG GCT) primers, in reactions with a final volume of 25 µL containing 2.5 µL of buffer solution, 2.0 µL of MgCl₂, 0.5 µL of DNTp, 0.2 µL of Taq DNA polymerase, 0.5 µL of each primer, and 1 µL of the sample. A PCR reaction occurred in a thermal cycler under the following conditions: 95 ºC for 2 min once (95 ºC for 30 min, 62.5 ºC for 30 min, 72 ºC for 1 min, and 72 ºC for 10 min 35x) and 6 ºC ∞ once. DNA was quantified with 1 µL of the product by agarose gel electrophoresis (0.8%). The PCR product was used for purification with the Illustra GFX PCR DNA kit and Gel Band Purification. The purification product was sequenced in duplicates using the Sanger method in an Applied Biosystems® ABI 3130 sequencer at the Center for Functional Genomics located at the Laboratory of Animal Biotechnology of the Luiz de Queiroz School of Agriculture. The 16S sequences were compared with the GenBank database (http://www.ncbi.nlm.nih.gov) with sequences known by similarity search via Blastn.

The similarity tree for bacterial isolate identification was built with MEGA 7 software (TAMURA et al., 2013). The Neighbor-Joining algorithm was used with the distance option according to the model proposed by JUKES & CANTOR (1969), with bacterium sequences being compared to type sequences based on the results of Blastn and the Ribosomal Database Project. The robustness of the similarity tree was confirmed by the analysis of 5,000 bootstrap replicates, and a Burkholderia sp. (AB334766-1) was used as the outgroup.

Microorganism testing for phosphate solubilization

Pure bacterial cultures isolated from the soil were inoculated in GL culture medium with 10 g of glucose and 2 g of yeast extract in 1 L of distilled water in test tubes, each containing 10 mL of the medium (SYLVESTER-BRADLEY, 1982). Tubes were maintained on an orbital agitation shaker (110 rpm, 28 ºC) for 24 h to obtain the pre-inoculum. A aliquot of 1 mL was used for optical density readings (OD600) in a spectrophotometer.

All bacteria had OD600 (0.5) equated by means of saline dilution (0.85%). Insoluble phosphate sources were separately added to each medium; 5 g L⁻¹ calcium phosphate [Ca₃(PO₄)₂], 1 g L⁻¹ iron phosphate (FePO₄), and 2 g L⁻¹ aluminum phosphate (AlPO₄). Then, 1 mL of each culture was inoculated in 8 mL of medium in triplicate (GADAGI...
& SÁ, 2002) and kept under constant agitation on a pendulum shaking table (TE-240/1, Tecnal®) at 100 rpm for 72 h at 28 °C.

After the growth period, the pH of the samples was measured and the phosphate solubilization was evaluated using the colorimetric method, in which 2 mL of the culture was centrifuged at 8,000 rpm for 10 min at 10 °C. Then, 1 mL of the supernatant was added to 9 mL of the working reagent [900 mL distilled water, 0.4 g ascorbic acid, and 100 mL 725 solution (0.1 g bismuth sub-carbonate, 6.8 mL sulfuric acid, 30 mL distilled water, 2.0 g ammonium molybdate, and 6.8 mL sulfuric acid for 100 mL in a volumetric flask)]. After 20 min, spectrophotometer readings were performed at 725 nm. Phosphate solubilization was determined using the phosphoric acid standard curve equation (160 mg L⁻¹).

Phosphate solubilizer microorganism growth

The growth curve of the microorganisms selected from phosphate solubilization was determined by inoculating a colony of each bacterium in test tubes maintained in an orbital agitation shaker (110 rpm, 28 °C) for 24 h. The measured OD600 was 0.6. Erlenmeyer flasks containing 40 mL of broth nutrient (5 g of peptone and 3 g of yeast extract in 1 L of distilled water) were inoculated with 400 µl in three repetitions. The OD600 was measured every 2 h until reaching the stationary phase. CFUs were counted by sample inoculation from plating in different rich media, considering the beginning of the growth and final log phase.

Results were subjected to analysis of variance and the means to Tukey’s test (p<0.05) using Sisvar® software. Pearson’s correlation was used for phosphate solubilization tests between soluble phosphate released by bacteria and the medium pH (p<0.05).

RESULTS AND DISCUSSION

Isolation, DNA extraction, and identification

CFU count was possible at 10² dilution. Only the minimal medium without herbicide treatment differed from the others (Figure 1). There were visual morphological differences in microorganism growth, and the rich medium treatments presented more accelerated growth, forming larger colonies in smaller quantities. Minimum medium treatments presented greater CFU formation, resulting in colonies of smaller diameter compared to those in the rich medium.

CFU growth was inhibited in the presence of herbicides in rich and minimal medium. This

![Figure 1 - Colony forming units of phosphate solubilizing microorganisms in rich medium with herbicide (RMWH); rich medium with no herbicide (RMWNH); minimum medium with herbicide (MMWH), and minimum medium with no herbicide (MMWNH). Histograms containing the same letters are statistically equal with reference to Tukey’s test (p<0.05).](image-url)
growth behavior could be related to the generation of reactive oxygen species (ROS) in cells, which increases upon exposure to chemical agents responsible for oxidative damage to cellular tissue (CRAY et al., 2013). Antioxidant enzymes present in microorganisms can be a defense against excessive ROS (GRATA et al., 2005). However, CFUs formed in the presence of the herbicide showed tolerance in rich medium. In the minimum medium, the strains used the herbicide as a nutrient or carbon source.

From the treatments with the herbicide, six CFUs were selected; two from rich medium, potentially tolerant, and four from minimum medium, potentially degrading. The selection parameter was the observation of morphological differences between CFUs grown on plates. The primary isolate identification involved the origin and growth medium used for the microorganism, with SCR-1 and SCR-2 in rich medium and SCM-3, SCM-4, SCM-5A, and SCM-5B in minimum medium.

The identification of the 16S rDNA sequence by similarity analysis compared with the GenBank database (http://www.ncbi.nlm.nih.gov) first identified six strains isolated from the soil, SCR-1, SCR-2, SCM-3, SCM-4, SCM-5A, and SCM-5B as Microbacterium arborescens, Bacillus pumilus, Stenotrophomonas maltophilia, Bacillus cereus, M. arborescens, and Bacillus safensis, respectively, with 100% similarity. The construction of the phylogenetic relationship identified five species (Figure 2). However, it was not possible to insert the species and SCM-5B due to the lower quality of the base pair sequence. However, similarity analysis compared with the GenBank estimated it as B. safensis.

Bacteria of the genus Bacillus can degrade a wide variety of xenobiotics (PINTO et al., 2012). When isolated from soil contaminated with herbicides, tolerance and degradation abilities were observed in tests with several molecules (ZHANG et al., 2018; CUI et al., 2018). Bacillus sp. showed high efficiency in atrazine removal (90%) in liquid media (GEED et al., 2017), in addition to increasing plant growth. The genus Microbacterium was investigated for environmental contaminant degradation and detoxification potential (AVRAMOV et al., 2016).

Phosphate solubilization

Tricalcium phosphate was the most efficient functional feature of solubilization among the tested sources. The maximum solubilization activity of \( \text{Ca}_3(\text{PO}_4)_2 \) was reached by the isolates B. pumilus (SCR-2) and S. maltophilia (SCM-3) with 334.44 g mL\(^{-1}\) and 334.30 g mL\(^{-1}\), respectively, with acidification of the pH of the culture medium for both strains, which favored the solubilization effect (Table 1).

A strong, negative \((r=-0.95)\) and significant \((p<0.05)\) correlation was observed

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Figure 2: Similarity analysis based on partial 16S rRNA sequences showing the relationship between bacterial isolates from contaminated soil and some of their closest phylogenetic relatives (indicated by the accession number). Burkholderia cepacia (AB3347661) was used as an external group.
between the concentration of solubilized tricalcium phosphate and pH, and the more acidic the pH of the culture medium, the more the inorganic phosphate is solubilized. MERBACH et al. (2009) observed that the calcium phosphate solubilization increases as the pH decreases. Organic acids produced by microorganisms can facilitate P solubilization by changing the pH of the medium (YADAV et al., 2017).

The FePO$_4$ solubilization showed effects on its concentration and the pH range, with a higher soluble phosphate concentration for the strains B. pumilus (SCR-2), with 271.33 mg mL$^{-1}$, followed by B. cereus (SCM-4), with 258.57 mg mL$^{-1}$ (Table 1). AlPO$_4$ showed no solubilization for the evaluated isolates.

The treatments control, M. arborescens SCR-1, B. pumilus (SCR-2), and B. cereus (SCM-4) showed a pH acidification considering that the culture medium with FePO$_4$ has an initial pH value of 6.5. However, the treatments S. maltophilia (SCM-3), M. arborescens (SCM-5A), and B. safensis SCM-5B presented basicity of the medium. The variables insoluble iron phosphate concentration and pH range showed a negative (r=-0.97) and significant (p<0.05) Pearson correlation coefficient. Therefore, iron phosphate is more solubilized as the pH of the medium is acidified. In this sense, the strains B. pumilus (SCR-2), with a pH of 4.53, and B. cereus (SCM-4), with a pH of 4.83, had the highest values of solubilized phosphate.

ANZUAY et al. (2017) studied bacteria of the genus Bacillus and obtained good FePO$_4$ solubilization and a decrease in the pH of the medium, followed by an increase in biomass through its inoculation in peanut and corn plants. Moreover, RAMANI (2011) observed an increase in phosphate solubilization in Bacillus strains even in the presence of herbicides.

AWAIS et al. (2017) carried out a study with the strain S. Stenotrophomonas maltophilia, isolated from sugarcane rhizosphere, and obtained significant values for the solubilization of calcium phosphates. According to ANZUAY et al. (2015), the amount of phosphate solubilizing bacteria does not change with an increase in pesticides, as they observed an increase in bacteria with this functional trait after the addition of herbicides and insecticides. According to the authors, the microorganism uses these molecules as a source of energy, carbon, and other nutrients in the cellular metabolism (DAS & DABNATH, 2006).

The strains B. pumilus (SCR-2) and S. maltophilia (SEM-3) were tolerant to hexazinone, with a functional trace of tricalcium phosphate solubilization, an important characteristic for soils with a high calcium content. According to KWON et al. (2017), Stenotrophomonas has biotechnological potential due to its metabolism. Also, B. pumilus was cited for having the ability to degrade organic molecules (COSTA et al., 2014) and characteristics for promoting plant growth (MEDINA et al., 2003). Phosphate solubilizing microorganism growth Measurements made at different intervals showed that for B. pumilus, the log phase began after 5 h of growth, the stationary phase was reached after 15 h, and there was a decrease after 18 h, as shown in figure 3. ANWAR et al. (2009) reported that B. pumilus presented brief insecticide degradation with chlorpyrifos at a high cellular density ($10^9$ CFU mL$^{-1}$). However, in lower densities, the degradation was slower, with a pH condition close...
to neutrality. AHMAD et al. (2012) observed the survival of *B. pumilus* in the rhizosphere of ryegrass after inoculation, with increased root area in the presence of the microorganism.

For the *S. maltophilia* bacterial strain, the log phase was observed after 5 h of growth and the stationary phase after 14 h, with decreased growth after 21 h (Figure 4). The genus *Stenotrophomonas* was reported
as a plant growth promoter and biological control agent due to its high metabolite and antifungal enzyme production (ELHALAG et al., 2015). For DUBEY et al. (2012), *S. maltophilia* has promising applications in bio- and phytoremediation due to genes that metabolize large varieties of organic compounds present in the rhizosphere of plants. It presents efficient xenobiotic (ZHANG et al., 2007), chlorpyrifos (DUBEY et al., 2012), and polyaromatic hydrocarbon degradation (TIWARI et al., 2016; SHUONA et al., 2017).

*S. maltophilia* can reduce MTBE concentrations (product added to gasoline) in water, and cellular growth increases until 18 h of incubation. Due to the characteristic genes of the bacteria, which can use the contaminant as the only carbon source, they may have increased growth with a degradation efficiency of 48% (ALFONSO-GORDILLO et al., 2016).

**CONCLUSION**

The isolation of bacteria from soil with a history of use of the herbicide hexazinone allowed selecting and identifying six bacterial isolates with potential for herbicide degradation. Among them, the strains *B. pumilus* (SCR2) and *S. maltophilia* (SCM3) stood out for the solubilization of tricalcium and iron phosphates. This research enables a sustainable alternative in agriculture with the use of these strains as soil bioremediators and phosphorus solubilizers, which are important factors for the increase in soil quality and promotion of plant growth.

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**DECLARATION OF CONFLICT OF INTERESTS**

The authors declare no conflict of interest. The founding sponsors had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, and in the decision to publish the results.

**AUTHORS’ CONTRIBUTIONS**

All authors equally contributed to manuscript design and writing.

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