Long Term Comparison of Talc- and Peat-Based Phytobeneficial 
*Pseudomonas fluorescens* and 
*Pseudomonas synxantha* 
Bioformulations for Promoting Plant Growth

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Use of plant growth promoting rhizobacteria (PGPR) is an important strategy in sustainable agriculture. Among PGPR, many *Pseudomonas* strains are of great interest due to their abilities to colonize and thrive in the rhizosphere, in addition to displaying mechanisms in plant growth promotion and biocontrol activities. However, as *Pseudomonas* strains are non-spore forming micro-organisms, their development into stable bioformulations for commercial applications can be difficult. This study examined over a long term period the effect of two different carriers, peat and talc, to prepare bioformulations using phytobeneficial *Pseudomonas* strains belonging to two taxonomical groups of interest: *P. fluorescens* and *P. synxantha*. Each strain has previously demonstrated plant growth promotion activity when inoculated in the rhizosphere. Each bioformulation was stored at ambient temperature and their viability was measured up to 180 days. In parallel, every 30 days (up to 180 days) each bioformulation was also applied in the rhizosphere of plantlets to validate their plant growth promotion activity, and their establishment in the rhizosphere was quantified by using strain-specific quantitative polymerase chain reaction assays. The viability of both *Pseudomonas* strains in the bioformulations was found to decrease after the first 15 days and remained relatively stable for up to 180 days. When applying the bioformulations to *Buglossoides arvensis* plantlets, the expected plant growth promotion was observed when using up to 180 day-old formulations of *P. fluorescens* and up to 120 day-old formulations of *P. synxantha*, with similar results for both carriers. Establishment of both *Pseudomonas* strains in the rhizosphere inoculated with the peat-based carrier bioformulations stored for up to 180 days was found to be stable. While a lower establishment of *P. fluorescens* in the rhizosphere was observed when talc-based bioformulations were stored for 90 days or more, rhizosphere colonization by *P. synxantha* talc-based bioformulations remained stable for up to 180 days. In conclusion, both peat and talc appear to be suitable carriers for *Pseudomonas* bioformulations,
INTRODUCTION

The world population is increasing at an alarming rate and is set to reach 9 billion by the year 2050, while the land available for agriculture has been decreasing due to industrialization (Aamir et al., 2020). Food production has increased through advances in agriculture such as the use of chemical pesticides and fertilizers, as well as breeding efforts to develop plant cultivars that are more productive and more stress tolerant (Aamir et al., 2020). However, plant productivity is still impacted by different environmental stresses such as drought, salinity, and limited nutrient availability (Basu et al., 2017; Aamir et al., 2020). Additionally, global climate change will also increasingly affect agriculture worldwide through increased surface temperature and reduced soil moisture (Basu et al., 2017). Conventional agriculture relies heavily on chemical pesticides and fertilizers to control pests and improve plant growth. The use of chemical pesticides varies among crops but data has shown that up to 93% of row crops such as corn, soybean, and cotton are treated with some kind of pesticide (Pimentel, 1993). Chemical pesticides have proven successful at preventing some crop diseases and enhancing crop productivity, however their role in damaging agro-ecosystems is now better understood (Mishra and Arora, 2016). Studies show that most chemical pesticides used in conventional agriculture can have detrimental effects on human health (Moses et al., 1993; Reigart and Roberts, 2013). In addition to their negative effects on human health, chemical pesticides can also negatively affect aquatic systems, the wildlife and the richness of soil microorganisms (Dorigo et al., 2009). The use of chemical fertilizers has also been beneficial to achieve greater crop productivity, however long-term experiments on the effects of fertilizers on soil fertility have shown that continual use can decrease soil quality (Kumar and Yadav, 2001; Yang, 2006).

Sustainable agriculture has been increasing in interest as a method to produce food to feed the growing population using environmentally friendly approaches. One attractive and economical approach is the use of plant growth promoting rhizobacteria (PGPR) to improve crop productivity and achieve higher yields (Antoun and Kloeper, 2001). PGPR are a functional group of bacteria that include different genera such as Rhizobium, Bradyrhizobium, Mesorhizobium, Azotobacter, Enterobacter, Xanthomonas, Bacillus, and Pseudomonas (Bashan, 1998; Bishnoi, 2015; Zayed, 2016; Berninger et al., 2018; Aamir et al., 2020). PGPR make up from 2 to 5% of all bacteria found in the rhizosphere (the soil area surrounding plant roots) (Antoun and Kloeper, 2001) and specific PGPR strains can be used alone or in combination to achieve different ecological functions leading to improved plant yields. Their use coupled with other crop management practices can lead to a reduction in the amount of chemicals used in the field without reducing crop yields. Among PGPR of interest, Bacillus and Pseudomonas are the most widely reported genera (Podile and Kishore, 2007). PGPR, and especially Bacillus and Pseudomonas, are of interest due to their ease of growth in the laboratory and their abilities to increase plant yields and/or improve plant health through various direct and indirect mechanisms of action. Direct methods include phosphate solubilization, nitrogen fixation, phytohormone production, iron acquisition, and the stimulation of plant defenses. Indirect methods include antibiosis, production of protective enzymes, induced systemic resistance, and the production of ACC-deaminase, which reduces the amounts of ethylene in the plant (Podile and Kishore, 2007; Martinez-Viveros et al., 2010; Zayed, 2016; Berninger et al., 2018; Bhattacharyya et al., 2020). Other beneficial traits of PGPR include their ability to efficiently colonize the root surface or the soil surrounding the roots, survive, compete and multiply in the soil (Bhattacharyya et al., 2017). PGPR that facilitate nutrient acquisition are known as biofertilizers, while PGPR that produce plant hormones are known as phytostimulators, and those that suppress diseases are known as biocontrol agents (Martínez-Viveros et al., 2010; Bishnoi, 2013). Some PGPR strains belong to more than one category. Biofertilizers, phytostimulators and biocontrol agents are considered key components of sustainable agriculture and are known to improve crop productivity in an ecofriendly manner. The biofertilizer market (comprising biofertilizers, phytostimulators, and biocontrol agents) is estimated to be worth 1.66 billion USD in 2022 and displayed an annual growth rate of 13.2% between 2015 and 2022, representing ~5% of the overall chemical-based fertilizers market (Macik et al., 2020).

A promising microbial inoculant must first be developed into a bioformulation to be used as a commercial product. A bioformulation is developed by combining a biologically active ingredient (living microbe or spore) with inert substances, known as carriers, to obtain a formulated product (Aamir et al., 2020). Common carriers include peat, talc, vermiculite, and diatomaceous earth (Bashan, 1998; Zayed, 2016). The carrier provides a temporary protective surface to the microorganism and must demonstrate the ability to support the growth of the target organism and maintain desired populations over an acceptable period of time (Macik et al., 2020). Carriers must have high water holding capacity, be well-buffered, environmentally safe, easy to use, and cost-effective (Bashan, 1998; Zayed, 2016; Bhattacharyya et al., 2020). Bioformulations are often divided into categories based on the composition of the carrier. These can include peat, liquid, granules, and freeze-dried powders (Abadias et al., 2005; Guijarro et al., 2007; Bhattacharyya et al., 2020). Among these, peat is one of the most common carriers in the biofertilizers market. Biofertilizers are often combined with other crop management practices to achieve different ecological functions leading to improved plant yields.

However, strain-specific variability exists and therefore the viability of each Pseudomonas strain and its capacity to maintain its plant growth promotion activity should be validated in different substrates before determining which formulation to use.

Keywords: bioformulation, Pseudomonas, peat, talc, carrier, plant-growth-promotion
carriers used worldwide (Bhattacharyya et al., 2020). Peat has a high water-holding capacity, high organic matter content, and is easily available (Bhattacharyya et al., 2020). Sticking agents are often added to peat-based biofertilizers to evenly distribute and stabilize the microorganism and can include carboxymethylcellulose, arabic gum, and polyalcohol derivatives (Bhattacharyya et al., 2020). Liquid formulations are aqueous suspensions made in oil, water, or a combination of both (Schisler et al., 2004). They are also a popular type of bioformulation; however, they can have reduced survival and propagation rates when compared to solid carriers (Berninger et al., 2018).

Developing a successful bioformulation can be a difficult process, as extensive trials must be conducted before the product can be marketed. Many bacteria are tested in pot assays under strict controlled settings, and their subsequent testing under field conditions can be inconsistent with pot studies or even unsuccessful (Lugtenberg and Kamilova, 2009; Backer et al., 2018). Many uncontrollable factors are found in the field such as variations between different field sites, year to year environmental variation, and even from growing different crops in the same field (Raaijmakers et al., 2009; Compan et al., 2010; Bishnoi, 2015). Soil is notoriously heterogeneous and introduced microorganisms can be unable to find an empty niche (Bishnoi, 2015). It is of utmost importance that the inoculated microorganism is able to compete with indigenous microorganisms and establish itself and survive in the rhizosphere. Many studies have shown that following a PGPR application, a sharp decline in its population is observed after a few weeks (Verma et al., 2013; Bishnoi, 2015; Corrêa et al., 2015; Novinscak and Filion, 2019). Additionally, the microorganism used to develop the bioformulation should be compatible with mass multiplication and not display any toxico logical effects to the plants or other beneficial microorganisms (Bishnoi, 2015). Another difficulty encountered when developing bacterial bioformulations is the use of Gram-negative strains because they do not produce spores (Berninger et al., 2018), which are often used for Gram-positive bacterial formulations. Gram-negative bacteria have however been found to be important PGPR, but due to their inability to produce spores, have a shorter shelf-life and are easily killed during desiccation of the formulation (Ramakrishna et al., 2019). One method used to counter the short shelf-life of Gram-negative bacterial formulations is to increase the number of microbes in the inoculant (Bashan et al., 2014; Tabassum et al., 2017). Even with a declining population over time, sufficient cells often remain alive to achieve a positive effect on the plants when applying the bioformulation in the field (Bashan et al., 2014).

_Pseudomonas_ strains are some of the most recognized strains of PGPR found in the rhizosphere (Podile and Kishore, 2007), however, they have not been widely commercialized in large part due to the rapid loss of viability of the available bioformulations (Corrêa et al., 2015). _Pseudomonas_ strains represent only 6% of the commercial biocontrol and biofertilizer products available for plants as compared to _Bacillus_ strains which represent 15% and _Trichoderma_ fungal species, which lead the way with 43% (Bettiol et al., 2012). _Pseudomonas_ strains are also found to be highly susceptible to environmental stress factors, such as drought and high temperatures (Paulitz and Bélanger, 2001). Greater commercial use of _Pseudomonas_ strains in crops will depend on the formulation of products with greater shelf life and the demonstration that the products are viable and effective when applied to plants. High moisture levels, favorable pH values and low temperature favors the survival of _Pseudomonas_ strains in carrier materials (Corrêa et al., 2015). Our research group has thoroughly examined the plant growth promoting effects of two strains of _Pseudomonas_ species, _Pseudomonas fluorescens_ LBUM677 and _Pseudomonas synxantha_ LBUM223. Previous studies conducted with LBUM677 have shown that in addition to plant growth promotion properties, the bacterium is able to increase seed oil content in various oilseed plants, including in corn gromwell (_Buglossoides arvensis_), a plant of high nutraceutical interest, under various field settings (Novinscak and Filion, 2018, 2019; Jiménez et al., 2020). LBUM223 has demonstrated plant growth promotion properties and has been successful in reducing the symptoms of the bacterial pathogen _Streptomyces scabies_ on potato plants in controlled and field settings (St-Onge et al., 2010, 2011; Arsenault et al., 2015). To reach the goal of eventually commercializing these strains, we wished to evaluate the survival and viability of these strains over a long period of time when produced in bioformulations. The present study examined their viability in two solid carriers, peat and talc, their capacity to colonize the rhizosphere when formulated in these carriers and stored for up to 180 days and finally their ability to conserve their plant growth promotion activity. Interestingly, many _Pseudomonas_ strains isolated to date which display strong PGPR abilities belong to the _P. fluorescens_ and _P. synxantha_ species. We therefore anticipate that the findings obtained in this study using LBUM677 and LBUM223 will, at least in part, be applicable to other _P. fluorescens_ and _P. synxantha_ strains of interest.

**MATERIALS AND METHODS**

**Bacterial Preparation**

_P. fluorescens_ LBUM677 and _P. synxantha_ LBUM223 were originally isolated from the rhizosphere of strawberry plants cultivated in Bouctouche, NB, Canada. Three replicates of pre-cultures of each bacterium were prepared by inoculating each bacterium in King’s B broth (King et al., 1954) and incubating at 25°C with agitation at 150 rpm for 24 h. Each pre-culture was used to inoculate larger volumes of King’s B broth (used for inoculum preparation) and incubated at 25°C with agitation at 150 rpm for 48 h. The bacterial concentrations were determined to range from 2.4 × 10^{11} to 3.7 × 10^{11} cfu mL^{-1} for LBUM677 and 8.8 × 10^{11} to 1.0 × 10^{12} cfu mL^{-1} for LBUM223 based on OD_{600nm} measurements and previously determined growth curves (Novinscak and Filion, 2018).

**Inoculum Preparation**

Talc (Fisher Scientific, Ottawa, Canada) and peat-based (Green Island peat moss, Isle-Verte, Canada) formulations of each _Pseudomonas_ strain were prepared by following the method of Vidhyasekaran and Muthamilan (1995). One large batch of each carrier was prepared by mixing eight kilograms of each carrier
with 80 g of carboxymethylcellulose (Sigma Aldrich, Oakville, Canada) and the pH was adjusted to 7 using calcium carbonate (Sigma Aldrich, Oakville, Canada). After adjusting the pH, eight aliquots of one kg of each substrate were prepared and were sterilized by autoclaving the carriers for 60 min at 121°C twice, on separate days.

Triplicates of each bacterium were inoculated in each carrier by aseptically adding 400 ml of bacterial suspension (see bacterial preparation) to 1 kg of carrier and mixing well. Duplicate negative controls were also prepared by aseptically adding 400 ml of uninoculated King’s B broth to 1 kg of carrier and mixing well. Each bioformulation (eight for each carrier type; three inoculated with LBUM223, three inoculated with LBUM677 and two negative controls, inoculated with King’s B broth) were air dried overnight and were then stored in the dark in polythene bags at room temperature (∼25°C).

Evaluation of the Shelf Life of the Talc and Peat Formulations

The viability of the bioformulations was tested after their storage for specific intervals of time (stored for 15 days, stored for 30 days, stored for 45 days, stored for 60 days, stored for 90 days, stored for 120 days, stored for 150 days, and stored for 180 days). After each storage time, one gram of each of the bioformulations was added to nine ml of sterile water, mixed, and six serial dilutions (from 10⁻² to 10⁻⁷) were prepared. One hundred µl of each dilution was plated in triplicate on Petri containing King’s B medium. The Petri were incubated at 25°C for 48 h before determining the number of colony-forming units on each plate.

Growth Promotion of Corn Gromwell Using Talc and Peat Formulations

After the bioformulations were stored for 30 days, 60 days, 90 days, 120 days, 180 days, the plant growth promotion effect of the bioformulations was also measured by mixing one gram of each bioformulation to 50 g of non-sterile soil and adding three Buglossoides arvensis seeds to each pot (to account for low germination rates). After germination, only one plant per pot was kept. The non-sterile soil used was obtained from a field at the Senator Hervé J. Michaud Agriculture and Agri-Food Canada Research Farm (Bouctouche, NB, Canada) and was characterized as a Gleyed Podzolic Gray Luvisol (GLPZ.GL), per the Canadian Soil Classification System with 62% sand, 25% silt, 13% of clay, 2.6% organic matter and a pH of 5.2 (Research Council of Canada, 1987). DNA was extracted from 0.5 g of rhizosphere soil using the method described in Griffiths et al. (2008). Following DNA extraction, all samples were diluted 1:10 before being used in qPCR. qPCR standards for LBUM223 and LBUM677 were created as previously described (deCoste et al., 2011) using a diluted cloned PCR fragment obtained from each bacterium using specific PCR primers. PCR primers and a TaqMan probe targeting a specific DNA sequence in LBUM223 were developed previously in Arseneault et al. (2016) (223phzCfor: 5'-ATA GAT GGA ATG CCG GTG ATG-3'; 223phzCrev: 5'-GCC CTC CTC CCT TTT TTG TTT-3' and 223phzCprobe: 5'-FAM-CGACAA ACT CCA GTC AA-MGBNFQ-3') while PCR primers and a TaqMan probe targeting a specific DNA sequence in LBUM677 were developed previously in Novinscak and Filion (2019) (BOX18F 5'-GCC TGC ATC GCG GTG ATG-3'; BOX18R 5'-CCC AAT CGG GTG ATC ATT G-3' and BOX18probe 5'-FAM-ATC ATA GAC TCG GAA TTG T-MGBNFQ-3'). The qPCR reactions were performed in 20 µl and consisted of: 10 µl of 10X iTaq Mix (Bio-Rad Laboratories, Mississauga, Canada), 0.8 µl of 5 µM of each probe, 0.8 µl of 5 µM of each primer, 3.0 µl of DNA and 4.6 µl of DEPC-treated H₂O. The amplification protocol was performed in a 7500 Real Time PCR machine (Applied Biosystems, Mississauga, Canada) and consisted of 95°C for 3 min followed by 50 cycles of denaturation at 95°C for 15 s and annealing/extension at 60°C for 60 s. Data collection was performed at the end of the annealing/extension step.

Statistical Analyses

The data from the bioformulation viability assay was rank transformed before statistical analysis. Data was separated by bacterium and a two factor ANOVA (sampling date and carrier type as factors) was used to examine the effects on the bacterial populations. For the plant growth promotion assays, each time-point was examined separately. Data were rank transformed when necessary. A two-factor ANOVA was used to measure the effect of the carrier type and the bacterial inoculation (LBUM677, LBUM223 or control) on plant weight. Finally, for the rhizosphere competency assay, the results for each bacterium were analyzed separately. A two-factor ANOVA examining the effect of the sampling date and the carrier type was performed. P-values below 0.05 were considered significant. SAS software version 9.2 (SAS Institute Inc, NC, USA) was used for all statistical analyses.

RESULTS

Evaluation of the Shelf-LIFE of Talc and Peat Bioformulations of P. fluorescens LBUM677 and P. synxantha LBUM223

LBUM677’s survival rate was higher in the peat carrier than the talc carrier (Table 1), while no difference was observed between the two carriers for LBUM223 (Table 1). For both bacteria, the population levels decreased almost two log values (cfu g⁻¹) from

Rhizosphere Competence of P. synxantha LBUM223 and P. fluorescens LBUM677

As mentioned previously, on each sampling date, rhizosphere soil was harvested from each plant, flash frozen in liquid nitrogen and stored at −80°C. A total of 320 rhizosphere samples were collected by the end of the experiment. The rhizosphere soil was subsequently lyophilized using a lyophilizer (Thermo Fisher Scientific, Mississauga, Canada). DNA was extracted from 0.5 g of rhizosphere soil using the method described in Griffiths et al. (2008). Following DNA extraction, all samples were diluted 1:10 before being used in qPCR. qPCR standards for LBUM223 and LBUM677 were created as previously described (deCoste et al., 2011) using a diluted cloned PCR fragment obtained from each bacterium using specific PCR primers. PCR primers and a TaqMan probe targeting a specific DNA sequence in LBUM223 were developed previously in Arseneault et al. (2016) (223phzCfor: 5'-ATA GAT GGA ATG CCG GTG ATG-3'; 223phzCrev: 5'-GCC CTC CTC CCT TTT TTG TTT-3' and 223phzCprobe: 5'-FAM-CGACAA ACT CCA GTC AA-MGBNFQ-3') while PCR primers and a TaqMan probe targeting a specific DNA sequence in LBUM677 were developed previously in Novinscak and Filion (2019) (BOX18F 5'-GCC TGC ATC GCG GTG ATG-3'; BOX18R 5'-CCC AAT CGG GTG ATC ATT G-3' and BOX18probe 5'-FAM-ATC ATA GAC TCG GAA TTG T-MGBNFQ-3'). The qPCR reactions were performed in 20 µl and consisted of: 10 µl of 10X iTaq Mix (Bio-Rad Laboratories, Mississauga, Canada), 0.8 µl of 5 µM of each probe, 0.8 µl of 5 µM of each primer, 3.0 µl of DNA and 4.6 µl of DEPC-treated H₂O. The amplification protocol was performed in a 7500 Real Time PCR machine (Applied Biosystems, Mississauga, Canada) and consisted of 95°C for 3 min followed by 50 cycles of denaturation at 95°C for 15 s and annealing/extension at 60°C for 60 s. Data collection was performed at the end of the annealing/extension step.

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TABLE 1 | Shelf-life of talc and peat bioformulations of *P. fluorescens* LBUM677 and *P. synxantha* LBUM223 determined by plating dilutions of inoculum at each time point.

| Carrier formulation and treatment | 15       | 30       | 45       | 60       | 90       | 120      | 150      | 180      |
|----------------------------------|----------|----------|----------|----------|----------|----------|----------|----------|
| Peat + LBUM677                   | 634.0 ab  | 306.0 ab  | 293.0 ab  | 205.0 ab  | 392.0 abc | 11.0 cdef | 3.7 ef   | 3.0 f    |
| Talc + LBUM677                   | 455.0 ab  | 45.1 bcd  | 12.5 cdef | 25.2 cde  | 8.4 def   | 6.0 ef   | 8.2 def   | 23.0 bcdf|
| Peat + LBUM223                   | 1190.0 ab | 268.0 ab  | 294.0 ab  | 197.0 ab  | 48.2 bcd  | 7.3 cde  | 4.9 ef   | 5.2 def   |
| Talc + LBUM223                   | 476.0 ab  | 105.0 bc  | 5.1 cdef  | 27.1 bcd  | 3.21 ef   | 6.1 cdef | 3.4 ef   | 4.2 ef    |

*Mean of three replications. *Means followed by a common letter in a row are not significantly different (P = 0.05) according to Tukey’s a posteriori test. The initial bacterial concentrations inoculated in the substrates were $2.81 \times 10^7$ cfu g$^{-1}$ for LBUM677 and $9.60 \times 10^7$ cfu g$^{-1}$ for LBUM223.

the initial inoculation of the carriers to the first sampling date, 15 days later (Table 1). A difference between the population levels of LBUM677 between peat and talc was first observed at 45 days and again at 90 days, however this difference between carriers was not maintained over time. Similarly, a difference between the LBUM223 population levels of both carriers was also observed at 45 and 90 days, however, as with LBUM677, the difference was not maintained. Over the course of the experiment, the population levels decreased by four log values from $\sim 1 \times 10^{11}$ to $1 \times 10^7$ cfu g$^{-1}$ for both bacteria in both substrates.

Growth Promotion of Corn Gromwell by *P. fluorescens* LBUM677 and *P. synxantha* LBUM223

The effect of the bacterial treatment was found to significantly influence the plant weight at all time-points measured when compared to the controls having received the carriers only without the bacteria (Figure 1). However, a significant effect of the carrier was only observed sporadically when the carriers reached 60, 90, and 180 days of storage (Figure 1). The plant growth promotion effect was observed for both bacteria at most time points, however, a difference between the effects of the two bacteria was initially observed when the bioformulations were stored for 90 days and this trend remained to the end of the experiment. The LBUM677 talc bioformulation was found to increase the plant weight from 12 to 380% (as compared to controls) at all time-point analyzed. The LBUM677 peat bioformulation was found to increase the plant weight from 37 to 364% at all time-points except for the 30 days bioformulation, which actually decreased the plant weight by 8% as compared to the control. The LBUM223 talc bioformulation was found to increase the plant weight by 93 to 130%, but only when the bioformulation was between 60 and 120 days of age. This was also observed in the LBUM223 peat bioformulation, where the plant weight was increased by 67 to 139% during the same time-points. Both LBUM223 bioformulations had a negative effect on the plant weight at 30 days and 180 days (Figure 1).

Rhizosphere Competence of *P. fluorescens* LBUM677 and *P. synxantha* LBUM223

When examining the rhizosphere competency of both bacteria evaluated using specific qPCR assays, LBUM677 was found to establish itself with higher population levels in the rhizosphere of corn gromwell than LBUM223 (Table 2). Additionally, LBUM677 populations were found to be higher when peat was used as a carrier than talc. When using the peat carrier, the LBUM677 populations were found to be stable throughout the experiment, however there was a significant decrease in the LBUM677 populations when talc was instead used. For LBUM223, no difference was observed in the rhizosphere competence between talc or peat carriers. The LBUM223 populations were found to be relatively stable throughout the entire experiment with significantly lower population levels observed at 120 days, however, this difference was no longer observed at 180 days.

DISCUSSION

The overall results of this study show that bioformulations stable up to 180 days at room temperature were successfully developed for two distinct *Pseudomonas* species. Both *P. fluorescens* and *P. synxantha* remained viable, could colonize the rhizosphere, and achieve plant growth promotion. When trying to determine if peat or talc would be a better carrier in general for *Pseudomonas* strains, our results indicated that both carriers yielded similar results at most time-points for both bacteria and demonstrated a shelf-life up to 180 days. Cell viability at 180 days in peat was similar for LBUM677 and LBUM223 (3 $\times$ 10$^7$ and 5 $\times$ 10$^7$ cfu g$^{-1}$, respectively), while results for talc were better for LBUM677 than LBUM223 (2 $\times$ 10$^6$ and 4 $\times$ 10$^7$ cfu g$^{-1}$, respectively). These results differ from those obtained comparing talc and peat as carriers for *Pseudomonas fluorescens* strains Pf1 and Pf2 showing that talc formulations were viable after 180 days of storage while peat formulations were only effective for up to 120 days of storage (Vidhyasekaran et al., 1997). The survivability of *P. fluorescens* strains Pf1 and Pf2 in six different carriers was tested and talc was also found to be the best carrier and maintained the highest populations of *P. fluorescens* at the end of 6 months of storage at 1.8 $\times$ 10$^6$ cfu g$^{-1}$ carrier (Gade et al., 2014). However, other results obtained using peat-based carriers, similar to the ones obtained in our study, showed that *P. fluorescens* Pf1, Pf2, and Pf27 remained viable for up to 8 months at a population of 7 $\times$ 10$^6$ cfu g$^{-1}$ (Vidhyasekaran and Muthamilan, 1995), and *P. chlororaphis* PA23 successfully maintained a shelf life of more than 6 months in a peat carrier (Nakkeeran et al., 2004). The survival of *P. fluorescens* PfALR2...
in peat, talc, and lignite was measured up to 60 days and peat was found to be a better carrier for this bacterium than talc and lignite (populations of $4.4 \times 10^7$ cfu g$^{-1}$ for peat as compared to $2.9 \times 10^7$ cfu g$^{-1}$ for talc) (Rabindran and Vidhyasekaran, 1996). Another study also tested three different carrier materials (cocopeat, talc, and farmyard manure) for their ability to support the growth of two *Pseudomonas* sp. isolates (An-2-nali and Pn-2-kho) (Verma et al., 2013). Cocopeat had the longest shelf-life of up to 7 months with a population of $1 \times 10^4$ cfu g$^{-1}$ (An-2-nali strain), while the shortest shelf-life was observed in the
The initial quantity of bacteria used to prepare the inoculum in this study was higher than what has been previously reported in the literature, where concentrations ranged from $1 \times 10^5$ to $2.5 \times 10^6$ cfu ml$^{-1}$ (Rabindran and Vidhyasekaran, 1996; Shaharoona et al., 2007; Jorgani et al., 2011; Jambhulkar and Sharma, 2013; Verma et al., 2013; Gade et al., 2014; Dubey and Kumar, 2018). The higher quantity determined in our study was achieved using a starter culture of each bacterium to inoculate the larger volumes required for the inoculations of the carriers. Studies have expressed that one of the ways to circumvent the decline in bacterial populations in a carrier is to use higher initial levels of the bacteria, however this approach might not always be economically efficient (Berninger et al., 2018). Even using higher initial amounts of cells, the population levels of both bacteria in our study decreased by four log-values from $1 \times 10^{11}$ to $1 \times 10^7$ cfu g$^{-1}$ carrier. A previous study found a rapid decrease in population levels of two *P. chlororaphis* strains (63-28 and TX-1) from $5 \times 10^8$ bacterial cfu ml$^{-1}$ to $1 \times 10^5$ bacterial cfu ml$^{-1}$ in peat and t alc in as little as 15 h after inoculation (Corrêa et al., 2015). However, in that study, the pH of t alc or peat was not adjusted to a neutral value of 7 (it was maintained at 9 for t alc and 3 for peat), which could explain the rapid decrease in bacterial populations observed. As increasing initial bacterial populations in carriers does not always lead to longer survival, efforts should be made to optimize storage conditions for long-term bacterial survival, including determining optimal storage temperatures and water content (Berninger et al., 2018).

In addition to a long shelf-life, bioformulations must also maintain their effectiveness to colonize their target plants. Interestingly, when examining this factor, LBUM677 was able to maintain its positive effect of promoting plant growth at almost all time-points except the initial time-point of 30 days (Table 2). The effects of LBUM223 on plant growth promotion were not as significant as LBUM677 (as compared to control) and even displayed a negative effect on plant weight at 30 days and 180 days as compared to controls. Differences in the activity of the two strains under study is most likely due to differences in their biochemical and/or genetic characteristics. This has previously been observed in bentonite and talc formulations of *Pseudomonas* sp. and CKK-3 on cotton seedlings. Strain was found to be more effective than strain CKK-3, likely due to differences in characteristics of this isolate (Sobhan et al., 2011). Other studies have shown the positive effects of *Pseudomonas* sp. bioformulations on various plant species. Specifically, four talc formulations of *Pseudomonas* sp. isolates CHAO, EPI, KKM, and VPT4 were successful in improving millable sugarcane height under field conditions (Viswanathan and Samiyappan, 2008). Wheat yield was improved by 15–25% over controls when using *Pseudomonas moravensis* and *Bacillus cereus* bioformulations (Hassan and Bano, 2015). Seed treatment of wheat with *P. fluorescens* RRb-11 in t alc led to significantly increased plant growth parameters and even increased seed germination by 94% over controls (Jambhulkar and Sharma, 2013). *Pseudomonas fluorescens* bioformulations showed the highest effectiveness of 8 different bioformulations on the fresh weight of sugar beets (Jorgani et al., 2011). *Pseudomonas putida* PT inoculated on maize seeds significantly enhanced the fresh and dry weights of the plants at harvest time (Khashei et al., 2020). Finally, chickpea seeds treated with t alc or peat formulations of *Pseudomonas* spp. effectively controlled chickpea wilt disease in two field trials and increased the chickpea yield (Vidhyasekaran and Muthamilan, 1995).

In addition to the effect of the biofertilizers on the plant’s weight, we also examined the persistence of each bacteria in the rhizosphere of corn gromwell. The rhizosphere competency of a bacterial strain is important for its ability to be a successful PGPR (Compant et al., 2019; Rilling et al., 2019). Also, before a new PGPR strain is developed into a bioformulation and commercialized for field applications, it is essential for regulatory considerations to develop reliable bioassays capable of detecting and quantifying the strain of interest among the complex soil microbiome (Mathre et al., 1999). In our study, specific qPCR bioassays were used for both strains under study, which successfully allowed to detect and quantify both strains in rhizosphere soil. Surprisingly, only few studies to date examined the population levels of inoculated bacteria in the rhizosphere and instead relied on increases in plant biomass and other plant parameters to confirm successful PGPR activity (Rilling et al., 2019).

Another important consideration when testing a bioformulation’s viability and establishment in the rhizosphere is the capacity to translate results obtained in controlled conditions, often in pots, to the field. Various *Pseudomonas* strains in peat

### Table 2: Rhizosphere competence of LBUM677 and LBUM223 bioformulations as determined by measuring the populations of each bacterium in rhizosphere soil using real-time PCR when the bioformulations reached different ages.

| Carrier formulation and treatment | Population (10$^7$ cfu per g) of bacteria in the rhizosphere soil detected using real-time PCR at various bioformulation ages (days) |
|----------------------------------|----------------------------------------------------------------------------------------------------------------------------------|
| Peat + LBUM677                   | 29.40 ± 0.07 at 30, 7.94 ± 0.07 at 60, 16.00 ± 0.07 at 90, 3.40 ± 0.07 at 120, 12.00 ± 0.07 at 180 |
| Talc + LBUM677                   | 1.53 ± 0.07 at 30, 7.04 ± 0.07 at 60, 0.71 ± 0.07 at 90, 0.23 ± 0.07 at 120, 0.11 ± 0.07 at 180 |
| Peat + LBUM223                   | 0.04 ± 0.07 at 30, 0.12 ± 0.07 at 60, 0.23 ± 0.07 at 90, 0.01 ± 0.07 at 120, 0.11 ± 0.07 at 180 |
| Talc + LBUM223                   | 0.16 ± 0.07 at 30, 0.34 ± 0.07 at 60, 0.08 ± 0.07 at 90, 0.01 ± 0.07 at 120, 0.31 ± 0.07 at 180 |

*Means followed by the same letter in a row are not significantly different (P < 0.05) according to Tukey’s a posteriori test.*
carrier have been tested in pot and field trials. Most strains displayed plant yield increases as compared to controls, however the results for a given strain were not consistent between pot and field trials (Shaharooona et al., 2007; Tabassum et al., 2017). Weather conditions, soils, plants, and the indigenous microorganisms in fields are all variable, leading to no specific rules for optimizing the inoculation of biofertilizers in soil (Khare and Arora, 2015). The success of the biofertilizer in field soil is dependent on its survival and ability to function in this heterogeneous environment. Abiotic factors (e.g., soil texture, pH, temperature and moisture levels) in the field exert their effects on inoculant population dynamics by imposing various stresses on the microbial cells (Evans et al., 1993; Khare and Arora, 2015). In our study, non-sterile field soil was used when performing pot studies, to help account for these abiotic factors found in this type of soil. This approach has previously allowed us to minimize the differences often observed between pot and field results with various PGPR and plant systems (St-Onge et al., 2011; Novinscak and Filion, 2018; Jiménez et al., 2020). The next step to validate the results obtained in this study will be to perform similar inoculation experiments under field conditions and track the PGPR strains in soil using the qPCR bioassays already available for LBUM223 and LBUM677. In parallel, the efficacy of the same carriers and inoculation approach will be validated for other Pseudomonas strains in development in our laboratory.

In conclusion, despite the apparent difficulty to develop stable bioformulations for non-sporulating plant beneficial Pseudomonas PGPR, significant progress is being made. Peat and talc-based carriers show great potential to assure viability of Pseudomonas strains at room temperature for many months and retain their ability to colonize the rhizosphere and achieve plant beneficial activity. Despite strain-specific differences, which need to be validated for each Pseudomonas strain of interest, both carriers yield similar results with no clear superiority of one over the other. The development of stable bioformulations is required for the commercialization of these organisms to be successfully used for plant growth promotion and/or biocontrol of plant pathogens in sustainable farming practices.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article-supplementary material, further inquiries can be directed to the corresponding author/s.

AUTHOR CONTRIBUTIONS

AN and MF contributed to the conception, design of the study, contributed to manuscript revision, read, and approved the submitted version. AN was responsible for conducting the experiments, the analyses, and writing the manuscript.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest. A patent has been granted on *P. fluorescens* LBUM677 and its use to enhance total lipid and SDA yields in an oilseed crop (US patent 10,165,745).

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