Prototipo de formulación y atmósfera de empaque para la cepa antagonista Pseudomonas fluorescens Ps006

Formulation prototype and atmosphere packaging for the antagonistic strain Pseudomonas fluorescens PS006

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Resumen

El aislamiento Pseudomonas fluorescens Ps006 demostró alto potencial para ser usado como principio activo de un bioinsumo, por su capacidad para producir biosurfactantes, actividad solubilizadora de fósforo y antagonista ante diferentes fitopatógenos. Por tal razón, el presente trabajo tuvo como objetivos desarrollar y caracterizar un prototipo de formulación a base de P. fluorescens Ps006, estable bajo condiciones de almacenamiento. Inicialmente se caracterizó el principio activo y se seleccionaron los auxiliares de formulación compatibles con el mismo, evaluándose la estabilidad de su mezcla con tres soportes sólidos, a dos humedades diferentes (10\% y 20\%) durante tres meses de almacenamiento a temperaturas de 8, 18 y 28 ± 2 °C. El principio activo demostró actividad antagonista in vitro sobre cuatro fitopatógenos y la temperatura y la humedad afectaron su estabilidad durante el almacenamiento. A los prototipos de formulación más estables en cuanto a viabilidad y actividad biocontroladora se les evaluó su estabilidad en presencia y ausencia de oxígeno y de protectores de membrana. Se seleccionó el soporte S1 al 20\% de humedad mezclado con el principio activo sin adición de protectores de membrana y almacenado en presencia de oxígeno, por ser el tratamiento más estable durante seis meses de almacenamiento, con pérdidas de viabilidad inferiores al 5\%.

Palabras clave: control biológico, formulación, Pseudomonas spp, vida útil.

Abstract

The isolation Pseudomonas fluorescens PS006 demonstrated high potential to be used as an active ingredient of a bio-product, because its capacity to produce biosurfactants, its phosphorus solubilizing activity and its antagonistic activity over different phytopathogens. For this reason, the present work had as objectives to develop and characterize a formulation prototype based on P. fluorescens PS006, stable under storage conditions. Initially the active ingredient was characterize and compatible formulation auxiliaries, were selected evaluating the stability of its mixture with three solid diluents at two different moistures (10\% and 20\%) during three months of storage at temperatures of 8, 18 and 28 ± 2 °C. The active ingredient showed in vitro biocontrol activity over four phytopathogens and temperature and humidity affected its stability during storage. Stability of the most stable formulation prototypes in terms of viability and biocontrol activity was evaluated in presence and absence of oxygen and membrane protectors. Support S1 with 20\% of moisture mixed with the active ingredient without addition of membrane protectors and stored in presence of oxygen, was selected as the most stable treatment during six months of storage at three temperatures, with viability losses lower than 5\%.

Key words: biological control agent, biocontrol, formulation, Pseudomonas spp, shelf-life.

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Introduction

*Pseudomonas fluorescens* is a bacterium with high potential for the biological control of different phytopathogenic microorganisms due to its multiple mechanisms of action, such as inducing systemic resistance, production of siderophores, competition for space and nutrients, and the production of different metabolites. Out of these metabolites, pyoluteorin, 2,4-diacetylphloroglucinol (DAPG), pyrrolnitrin, hydrogen cyanide and other metalloenzymes are of note, (Loper & Gross, 2007; Anjaiah, 2006; Manikandan et al. 2010; Siddiqui & Shaukat, 2003). Some of the diseases over which *P. fluorescens* exercises control are sheath blight in rice (*Oryza sativa* L.) caused by *Rhizoctonia solani* (Commare et al. 2002; Rabindran & Vidhyasekaran, 1996); damping-off in tomatoes (*Solanum lycopersicum* L.) caused by *Fusarium oxysporum* (Manikandan et al. 2010; Valencia et al. 2005); bacterial spot in tomatoes caused by *Xanthomonas campestris* pv. *Vesicatoria* (Kamal et al. 2008); rotting of the fruit and mold in chilies caused by *Colletotrichum capsici* and *Leveillula taurica*, respectively (Anand et al. 2010); and wilt caused by the soil fungus *Verticillium dahlia*, a disease responsible for great economic losses in crops such as cotton (*Gossypium* sp. L) (Erdogan & Benlioglu, 2010).

Like other microorganisms, *P. fluorescens* is susceptible to different environmental conditions, particularly temperature and pH (O’Callaghan 2006). Therefore, for its efficient implementation as a biological control tool, it is necessary to develop products with a formulation that ensures their effectiveness and stability during storage (Burges, 1998).

There are many products around the world with *P. fluorescens* as the active ingredient that are formulated for the biological control of different plant pathogens (Hernández-Rodríguez et al. 2008; Hernández-Rodríguez et al. 2014). These products include BioCure-B®, recommended for the control of *Mycosphaerella grasicola*, *Pythium spp.*, *Rhizoctonia solani*, *Fusarium spp.*, *Botrytis cinere*, *Sclerotium rolfsii*, and *Sclerotinia homoeocarpa* (T-Stanes & Company Limited, s.f), and BlightBanaA506® formulated as a wettable powder for the control of *Erwinia amylovora* in potato (*Solanum tuberosum* L.) and tomato crops (Copping, 2009), among other products.

The autochthonous strain *P. fluorescens* Ps006 has the capacity of producing biosurfactants, a phosphorus solubilizing activity, and a zoosporicidal activity, as well as the potential for biological control of pathogens by showing a reduction in the severity index (SI) of *O. virulentus* (SI of 0.6) in lettuce plants (*Lactuca sativa* L.) compared to the control pathogen (SI of 4.7). Furthermore, this bacterium applied at a concentration of 1 x 10^7 cells per ml facilitated and significantly increased the length and width of the leaves, the dry biomass of the aerial part, as well as the development in the length and dry biomass of the root of fique plants, demonstrating a promoting effect on plant growth (Smith et al. 2013). Taking said characteristics into account, this research aimed to develop a prototype for the formulation of wettable powder (WP) using *P. fluorescens* Ps006 as an active ingredient, and to select the most suitable packaging atmosphere to maintain the viability of the microorganism.

Materials and Methods

**Microorganism and Conservation**

The isolation of *P. fluorescens* Ps006 was used, obtained from the rhizosphere of fique *Furcraea andina* (Trelease 1808) in the municipality of Totoró (2°38’ N and 2°15’ W, at 2,750 m.a.s.l and 14 °C), in the Cauca Department, Colombia (Sastoque, 2010). The microorganism was conserved in saline solution (0.85% NaCl) with 10% glycerol and 0.10% peptone at -70 ± 2 °C.

**Production and Characterization of the Active Ingredient of *P. fluorescens* Ps006**

*P. fluorescens* Ps006 was planted in Luria Bertani (LB) agar medium (Oxoid CM1021) and was incubated for 48 hours at 28 ± 2 °C. From this culture, a cell suspension was prepared in Tween 80 at 0.5%, and it was inoculated in a 250-ml Erlemmeyer flask with 100 ml of LB medium. The concentration of the suspension was estimated by reading the absorbency at a wavelength of 300 nm, and said value was extrapolated in a previously standardized calibration curve. Fermentation was carried out using constant agitation of 175 rpm at 28 ± 2 °C for 48 hours.

Three fermentation batches were developed, which were characterized by determining the cell concentration in colony-forming units per milliliter (CFU/ml), the pH and the *in vitro* biological activity, three times for each batch.

The bacterial concentration was determined through a count in a Petri dish, planting three decimal dilutions of the fermentation broth three times in Petri dishes with LB medium and incubating them for 24 hours at 28 ± 2 °C, when the CFU/ml count was carried out. Once the normality (Shapiro-Wilk test 95%) and homogeneity of variance (Bartlett's test 95%) of the data was verified, an analysis of variance (ANOVA) and Tukey's test were carried out with a confidence level of 95%. The Statistix 8.0 statistical program was used.

The pH of the three fermentation batches was measured with a previously calibrated Hanna® Instruments brand potentiometer. The standard deviation and the coefficient of variation of the data were determined.

The *in vitro* biological activity was estimated as the growth inhibition of four recognized plant pathogens (Raut et al. 2012) through the dual culture technique.
Results and Discussion

Production and Characterization of the Active Ingredient P. fluorescens Ps006

An average pH of 8.63 was obtained for the three active ingredient batches with a 0.2% coefficient of variation. The cell concentrations for each one of the evaluated batches were $6.73 \times 10^8$, $8.07 \times 10^8$ and $8.93 \times 10^8$ CFU/ml. The coefficient of variation of the evaluation results was 0.6%, and there were no significant differences between the results of the batches ($F = 2.06; DF = 2; p = 0.2081$), which indicates low variability and high repeatability in the production of the active ingredient (Villamizar et al. 2005). The pH oscillated between 8.60 and 8.65, finding no significant differences between the results of the evaluated batches ($F = 3.74; DF = 2; p = 0.0882$).

Based on the results obtained for each parameter, ranges or limits of acceptance were proposed in this study for the characteristics of the active ingredient based on Ps006, which will be used as a reference for its quality control (Table 1).
In Figure 1, the in vitro effect of the active ingredient on the four evaluated pathogens is observed with inhibitions of the diameter growth of R. solani Rh200, B. cinerea Bc008, S. sclerotiorum Sc021 and F. oxysporum MAP5 between 93.64% and 36.80%. The antagonistic activity of P. fluorescens toward pathogens that affect crops of economic importance has been demonstrated by several authors. For example, for F. oxysporum, a control of 81.21% was achieved in field conditions (Manikandan et al., 2010), and for R. solani, decreases of 42% were observed in the intensity of the disease (Rabindran & Vidhyasekaran, 1996) and of 47.89% (Commare et al., 2002) in greenhouse conditions. Likewise, a 67.7% decrease in the in vitro growth of Botrytis sp. was achieved (Mikani et al., 2008), as well as a 26.3% reduction in the incidence of the disease for S. sclerotiorum in field conditions (Fernando et al., 2007). These results confirm the potential of P. fluorescens as a biological control agent and demonstrate the potential of its use as an active ingredient in biopesticides for the control of pests that affect crops of economic importance. However, research on the scale of plant pots and fields is necessary to demonstrate this hypothesis.

The most resistant pathogen to the antagonistic activity of P. fluorescens Ps006 was F. oxysporum MAP5 with an inhibition of the diameter growth of only 36.80%, a significantly lower value (F = 14.9; DF = 3; p = 0.0000) to those obtained with the other evaluated pathogens, which presented inhibition percentages above 87%. This could be due to a greater resistance in the cell wall of the F. oxysporum mycelium; a microorganism that has polymers of heteroglucans that are not found in the other evaluated pathogens, which have been associated with the resistance and maintenance of cell integrity (Nuero, 1995). It is also possible that the isolation of F. oxysporum MAP5 has the capacity to inhibit the biosynthesis of 2,4-diacetylphloroglucinol (2,4-DAPG); an antibiotic with a wide spectrum that is recognized as one of the main mechanisms of action in a wide range of isolations of P. fluorescens (Showkat et al., 2012). This behavior was demonstrated by Schouten et al. (2004), who evaluated the sensitivity of 41 isolations of F. oxysporum to said antibiotic, and found resistant isolations of 17%, which attributed to the capacity of this pathogen to produce fusaric acid, a powerful inhibitor of the biosynthesis of 2,4-DAPG.

Table 1. Limits of acceptance established for the characteristics of the active ingredient.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Average Experimental Value</th>
<th>Limits</th>
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<tbody>
<tr>
<td>Concentration (CFU/ml)</td>
<td>7.9 x 10⁹</td>
<td>&gt; 1.0 x 10⁹</td>
</tr>
<tr>
<td>pH</td>
<td>8.63</td>
<td>8 - 9</td>
</tr>
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The mode of action of P. fluorescens as an antagonist of plant pathogens has been related to the competition for space and nutrients. This phenomenon was observed by Commare et al. (2002), who determined the antagonistic activity of formulations of P. fluorescens in talc, showing 47.89% control of R. solani in greenhouse conditions, the results being attributed to competition for space. In the study conducted by Valencia et al. (2005), it was concluded that the activity of P. fluorescens ZUM80 on F. oxysporum was due to competition for nutrients, depriving the fungus of the iron available in the medium through the production of siderophores. This antagonistic activity was facilitated when the bacteria had pre-exposure time in the medium (Valencia et al. 2005). Apart from these mechanisms of competition in studies conducted by Khanam, Ueno, Kihara, Honda & Arase (2005), it was found that the salicylic acid produced by P. fluorescens inhibits the formation of infection structures such as the germ tube and appressoria in B. cinerea, which suggests that this bacterium also has the capacity to produce other metabolites with effective antagonistic activity.

Selection of Compatible Solid Supports

The total loss of viability of the active ingredient after three months of storage at the three evaluated temperatures is shown in Figure 2. For the S1 support at 10% humidity (S1-10%), losses between 14% and 31.9% were observed, and for S1 at 20% humidity (S1-20%), losses between 9.6% and 33.2% were observed. For the S2 support at 10% humidity (S2-10%) at the three temperatures, a 67.7% loss of viability was observed, and for S2 at 20% humidity (S2-20%), the reduction in the viability oscillated between 2.3% and 9.4%. Finally, for the S3 support at 10% humidity (S3-10%) at the three temperatures, a 78.1% loss was observed, and for S3 at 20% humidity (S2-20%), the loss of viability oscillated between 25.7% and 69.6%.

![Figure 1. Antagonistic in vitro activity of the active ingredient based on P. fluorescens Ps006, determined as the diameter growth inhibition of four plant pathogenic fungi. Treatments with the same letters do not present significant differences according to Tukey’s range test (95%).](image-url)
The results showed that the 10% and 20% humidity conditions, to which the systems were adjusted for their storage, had an effect on the stability of the Ps006 bacterium’s viability. The loss of viability with S2 and S3 at 10% was significantly higher than for the systems with 20% humidity (F = 402; DF = 17; p = 0.0000) (Figure 2). This may be because the treatments with 20% humidity required the addition of a volume of buffer solution at pH 7.5 to adjust the humidity. This possibly exercised control on the ionic state of the formulation, making it less susceptible to abrupt changes in pH, which may affect the stability and integrity of the cell membrane, cause the denaturation of enzymes, and alter the ionic interactions that allow them to recognize and bind to the substrate (Calvo et al. 2004; Nelson & Cox, 2005).

The greatest losses in viability obtained with S2 and S3 when the humidity was adjusted to 10% could be because at said humidity, the concentration of solutes in the system is greater compared to the treatment at 20% humidity. This hypertonic atmosphere in the medium at 10% humidity could have caused plasmolysis phenomena, forcing the cell to lose water, which increases the viscosity of the intracellular environment, decreasing its volume, causing a retraction of the cell membrane, and subsequently, cell death (Rodríguez et al. 2005; Tortora et al. 2007).

With the S1 support at both humidities and with S2 and S3 at 20% humidity, it was observed that the loss of viability increased as the storage temperature increased, a negative effect that has been shown by different authors for diverse microorganisms (Chen et al. 2005; Nelson & Cox, 2005; Kinay and Yildiz 2008) evaluated formulations of *Pichia guilliermondii* and observed that those stored at 4 °C were more stable than those stored at 24 °C. Similar results to those presented in the present study were obtained by Santos et al. (2012), who evaluated the stability of germination of conidia in the Colombian isolations Trichoderma koningiosis Th003 and Trichoderma asperellum Th034. They were formulated as dry powders for dusting and dispersible granules stored at temperatures of 8, 18, and 28 °C. Said authors obtained longer useful lives when storage was at a temperature of 8 and 18 °C, concluding that the lowest temperatures reduced the metabolic activity of the conidia, preventing the production of toxic metabolites and the exhaustion of nutrients; aspects related to microbial physiology.

The active ingredient mixed with S1 and S2 at 20% humidity were selected as the basic delivery systems for the development of a biopesticide, because they were the most compatible systems with the active ingredient *P. fluorescens* Ps006, maintaining stable viability and biocontrol activity during three months of storage at 8 ± 2 °C, 18 ± 2 °C and 28 ± 2 °C.

**Evaluation of Adjuvants and Packaging Atmosphere**

Once the most compatible supports with the active ingredient had been selected, the effect of the presence and absence of oxygen in the packaging and of the two protective adjuvants of the cell membrane (skimmed milk and glycerol) on the viability of *P. fluorescens* Ps006 in storage conditions was evaluated. The results of said tests are presented in Figure 3.

Taking into account the concentration of the cell suspension (1 x 10⁸ CFU/ml) and the proportion of the mixture with the supports, the theoretical concentration of the mixture is estimated to be 1.3 x 10⁸ CFU/ml. The experiment data showed that the initial viability of the active ingredient with the S1 support at 20% humidity was 4.67 x 10⁷ CFU/g, and when it was mixed with the membrane protectors, it was 1.45 x 10⁸ CFU/g. This decrease in the cell concentration when a membrane protector was not used is possibly related to a toxic effect of the support by entering into contact with the *P. fluorescens* Ps006 cells, which reduced its viability. However, this deleterious effect was decreased by the membrane protectors.

It was observed that the loss of viability after six months of storage was directly proportional to the temperature, a trend that was evident in both atmospheric conditions (with and without oxygen). The treatment without a membrane protector and stored with oxygen was more stable for S1, because the maxi-
mum loss of viability was 20% at 28 °C, compared to the other treatments with significantly higher losses (F = 1,622.39; DF = 10; p = 0.0000), which exceeded 70% under the same temperature conditions.

For S2, initial viabilities of 1.79 x 10^7 CFU/g were obtained when membrane protectors were not used and of 6.90 x 10^7 CFU/g when they were added, again showing an initial toxic effect upon contact with the solid support. In general, after six months it was observed that the loss of viability increased as the storage temperature increased. Said loss was significantly lower (F = 233; DF = 9; p = 0.0000) for the treatments stored with oxygen compared to the treatments stored in a vacuum. This means that the absence of oxygen in the packaging did not improve the stability of the formulations. This behavior could be due to the stress caused by the vacuum atmosphere on an obligate aerobic bacterium such as *P. fluorescens*, which generates energy by a process of oxidative phosphorylation with oxygen as its last electron acceptor (Adams & Moss, 2008; Vásquez et al. 2009). The negative effect of the vacuum packaging on the viability of *Pseudomonas* spp. showed in this study has been widely studied in the food industry, where vacuum packaging is used to inhibit the development of contaminating bacteria (ICMSF, 2006). Preventing the need to implement vacuum systems is an evident advantage for industrial production, because the times and costs of the process are reduced, thus increasing the profitability of the end product.

In most of the cases, the viability was significantly higher (F = 233; DF = 9; p = 0.0000) for the treatments in which the active ingredient was only mixed with the support, compared to the treatments in which, as well as the support, the active ingredient was mixed with the potential membrane protectors. This was with the exception of the S2-20% treatment stored with oxygen, which did not show significant differences when the protectors were used or not. Therefore, glycerol is widely used as a humectant and osmotic protector (Freitas et al. 2009; Rowe et al. 2009), as well as a cryoprotectant and protector against dryness, such as skimmed milk (Cody et al. 2008; Morgan et al. 2006; Barbaree et al. 1982). However, it has been found that certain strains of *Pseudomonas* spp. use glycerol as a source of carbon to produce biosurfactants (Freitas et al. 2009; Freitas et al. 2010; Silva et al. 2010; Stanghellini & Miller, 1997). This could suggest possible consumption of glycerol and skimmed milk during storage by the formulation prototypes evaluated in the present study. This may have facilitated the growth of the microorganism, maintenance of the cell metabolism and the production of toxic metabolites, causing losses of viability (Santos et al. 2012).

The development of a powder formulation of antagonistic bacteria is especially important for the treatment of seeds (Ramamoorthy et al. 2001). The present study developed a formulation prototype based on the bacterium *P. fluorescens* Ps006, which survived during six months of storage at 28 °C, having a minimum loss of viability when it was formulated in S1, which is a clay adjusted to 20% humidity, and was used in an uncontrolled packaging atmosphere with air. Similar formulation prototypes have been developed in other studies, but with less stability and in the majority of cases, requiring refrigeration conditions. Such is the case of populations of *P. fluorescens* formulated in talc with 20% xanthan gum, which maintained their viability during two months of storage at 4 °C (Kloepper & Schroth, 1981). Another formulation that used a vermiculite clay like the present study obtained a stable viability for six months at 4 °C (Connick, 1988). In another study, where the support was talc, the viability was maintained for 90 days at 26 °C (Sadi & Masoud, 2012).

![Figure 3](image-url) **Figure 3.** Loss of viability of the active ingredient *P. fluorescens* Ps006 with the supports **A)** S1-20% and **B)** S2-20% after six months of storage. CV: Stored in a vacuum; CO: Stored with oxygen; PPV: Stored in a vacuum with membrane protectors; and PPO: Stored with oxygen with membrane protectors. Treatments with the same letter do not present significant differences according to Tukey’s test (95%).
Conclusions

The prototype for formulation in wettable powder (WP) with the active ingredient *P. fluorescens* Ps006 as a base, which demonstrated its antagonistic activity against the plant pathogens *Fusarium oxysporum*, *Rhizoctonia solani*, *Botrytis cinerea* and *Sclerotinia sclerotiorum*, was stable for six months of storage at the three evaluated temperatures, when clay adjusted to 20% humidity was used as a support without the addition of membrane protectors and using a packaging atmosphere with oxygen. The losses of viability were less than 5% during the time of storage without refrigeration. Therefore, this prototype was selected to continue with the development of a commercial product with the potential to control diseases in different crops.

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References


Formulation Prototype of *Pseudomonas fluorescens*