Response of the Biocontrol Agent *Pseudomonas pseudoalcaligenes* AVO110 to *Rosellinia necatrix* Exudate

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ABSTRACT

The rhizobacterium *Pseudomonas pseudoalcaligenes* AVO110, isolated by enrichment of competitive avocado root tip colonizers, controls avocado white root rot disease caused by *Rosellinia necatrix*. Here, we applied signature-tagged mutagenesis (STM) during the growth and survival of AVO110 in fungal exudate-containing medium with the goal of identifying molecular mechanisms linked to the interaction of this bacterium with *R. necatrix*. A total of 26 STM mutants out-competed by the parental strain in fungal exudate, but not in rich medium, were selected and named growth-attenuated mutants (GAM). Twenty-one genes were identified as required for this bacterial-fungal interaction, including membrane transporters, transcriptional regulators and genes related to the metabolism of hydrocarbons, amino acids, fatty acids and aromatic compounds. The bacterial traits identified here that are involved in the colonization of fungal hyphae include proteins involved in membrane maintenance (a dynamin-like protein and ColS) or cyclic-di-GMP signaling and chemotaxis. In addition, genes encoding a DNA helicase (*recB*) and a regulator of alginate production (*algQ*) were identified as required for efficient colonization of the avocado rhizosphere.

IMPORTANCE

Diseases associated with fungal root invasion cause significant losses on fruit tree production worldwide. The bacterium *Pseudomonas pseudoalcaligenes* AVO110 controls avocado white root rot disease caused by *Rosellinia necatrix* using mechanisms involving competition for nutrients and niches. Here, a functional genomics approach was conducted to identify bacterial traits involved in the interaction with this fungal pathogen. Our results contribute to a better understanding of the multitrophic
interactions established among bacterial biocontrol agents, the plant rhizosphere and the mycelia of soilborne pathogens.
INTRODUCTION

The significance of multitrophic interactions established among organisms during the biological control of soilborne plant pathogens has been extensively reported (1,2,3) and it is widely accepted that successful colonization of the plant rhizosphere is a relevant biocontrol trait (4). In this regard, many studies have focused on the identification of bacterial genes involved in the attachment and colonization of plant roots and seeds with special emphasis on biocontrol Pseudomonas (5,6,7,8,9). However, little attention has been given to the trophic and physical interactions established between bacterial biocontrol agents and root pathogenic fungi. Biocontrol Pseudomonas spp. have been shown to interact with fungal pathogens via chemotaxis and by living in close proximity to fungi, often colonizing hyphal surfaces and utilizing nutrients exuded from living fungal cells (1,10). In fact, this bacterial behavior has been proposed to contribute to biocontrol through the biosynthesis of antifungal compounds or the release of enzymes involved in the degradation or alteration of fungal components (11).

The biocontrol rhizobacterium Pseudomonas pseudoalcaligenes (Proteobacteria, Gammaproteobacteria, Pseudomonadales) strain AVO110, isolated by enrichment of competitive avocado (Persea americana) root tip colonizers, is able to control avocado white root rot disease caused by the soilborne pathogen Rosellinia necatrix (Ascomycota, Sordariomycetes, Xylariales) under greenhouse conditions, reducing disease development up to 45% when compared to control plants not inoculated with bacteria (12). Although most bacterial strains isolated using this strategy were shown to produce several exoenzymes, hydrogen cyanide (HCN) or antifungal antibiotics, the potential biocontrol traits of P. pseudoalcaligenes AVO110 only included the biosynthesis of siderophores, weak cellulose activity (12) and its ability to colonize both...
the avocado rhizosphere and the *R. necatrix* hyphae (1). Thus, competition for nutrients and niches was proposed as the most relevant biocontrol trait of this bacterium (1). Actively growing fungal hyphae exude a complex mixture of low molecular weight metabolites that include organic acids, such as oxalic, citric and acetic acid, peptides, amino acids, sugars and sugar alcohols, such as mannitol (13,14,15,16). Along this line, *P. pseudoalcaligenes* AVO110 was shown to efficiently grow on minimal BM medium only when supplemented with *R. necatrix* exudates (BM-RE medium), reaching higher cell densities than other non-biocontrol rhizobacterial strains (1). Thus, AVO110 might encode specific traits conferring a competitive advantage to this bacterium during its interaction with fungi.

Several strategies are currently available to unravel bacterial genes involved in host interactions, including transcriptional profiling, *in vivo* expression technology (IVET) and signature-tagged mutagenesis (STM) (17,18,19). However, few studies have addressed the identification of bacterial genes involved in interactions with fungal phytopathogens. During a commensal interaction, the plant pathogenic fungus *Gaeumannomyces graminis* was shown to induce *Pseudomonas fluorescens* Pf29Arp genes involved in mycelium colonization even before cell-to-cell contact (20). Likewise, in a non-contact confrontation assay, the mycophagous bacterium *Collimonas fungivorans* responded to *Aspergillus niger* by activating genes for the utilization of fungal-derived compounds and for production of a putative antifungal compound (21). In two studies that applied IVET, *P. putida* 06909 genes induced during the colonization of *Phytophthora* mycelia were identified (22, 23). This strategy yielded several genes involved in carbon and amino acids metabolism, ABC transporters, and outer membrane porins. Furthermore, colonization of *A. niger* hyphae by *B. subtilis* was suggested to be an active process in which the bacterium rewires not only surface...
attachment, but also metabolism, motility, general stress responses and antimicrobial production (24). In spite of all these exciting new insights into the interactions established between bacteria and fungi, genes revealed by transcriptional profiling or IVET should be further investigated experimentally by functional approaches. In this sense, STM combines the power of insertional mutagenesis and negative selection with a detection system, which allows the identification of individual attenuated mutants from a complex mutant pool. Previously, we showed that the utilization of fungal exudates plays an important role in the biocontrol ability of P. pseudoalcaligenes AVO110 against R. necatrix (1). In this study, we used this bacterial-fungal model system to apply STM in the identification of bacterial genes involved in interactions with a fungal phytopathogen. Sequence analysis of the genes interrupted by the transposon in the selected P. pseudoalcaligenes AVO110 mutants revealed several molecular processes involved in the interactions of this bacterium with R. necatrix, including genes related to the colonization of biological surfaces and utilization of fungal exudates.

RESULTS

Selection of Pseudomonas pseudoalcaligenes AVO110 Growth Attenuated Mutants in Rosellinia necatrix Exudates. A mutant bank of 3,408 P. pseudoalcaligenes AVO110 miniTn5Km2-tagged derivatives was constructed as described in Materials and Methods using the strains listed in Table 1. Grouped in 76 input pools of ≤ 45 mutants, all 3,408 STM mutants were screened for growth in minimal BM medium containing R. necatrix exudates (BM-RE media, pH=7.11). All input pools included a negative control (wild-type P. pseudoalcaligenes AVO110) and a positive control (a
miniTn5Km2-tagged AVO110 derivative selected by its ability to growth and survive on both LB and BM-RE). A total of 765 strains showing stronger hybridization signals with the input probe than with the output probe was selected for further characterization.

To reduce the number of false-positive candidates, the selected mutants were grouped in new pools, mixed with other random mutants, and re-tested in a second round of STM screening. After this second STM round, the number of mutants was reduced to 99 strains (Fig. S1).

Single insertion of the transposon into the genome of each mutant was determined by hybridization of EcoRI/BglII-digested total DNA against a transposon probe (aphA gene). Out of the 99 mutant strains, 85 contained a single insertion of the transposon and were selected for further analysis (see Materials and Methods). These strains are potentially mutated either in essential genes or in genomic regions required for competitive survival in BM-RE. To differentiate between these two possibilities, competitive indexes (CIs) of each of the selected mutants in comparison with the wild-type strain were calculated both in LB medium (CI_{LB}) and BM-RE (CI_{BM-RE}). A total of 26 mutants showed CI_{BM-RE} and CI_{LB} values significantly less than 1 and not significantly different from 1, respectively (CI_{BM-RE} < 1 and CI_{LB} = 1) (Fig. 1). Thus, these strains, which were named as growth attenuated mutants (GAM), were out-competed by the wild-type strain only in media supplemented with fungal exudates.

**Pseudomonas pseudoalcaligenes** AVO110 Genes Required for Growth in Fungal Exudates. Genomic DNA fragments flanking the transposon insertion within the 26 selected GAM strains were cloned, sequenced, and used to search the RAST-annotated draft genome sequence of *P. pseudoalcaligenes* AVO110 generated in this study (see Materials and Methods). Furthermore, the nucleotide sequences of the interrupted
AVO110 genomic regions were used to search the GenBank and ASAP databases. After identification of several independent insertions of the transposon in the same gene (colS, leuC and recB) and insertions in intergenic regions (strains GAM13 and GAM19), a total of 21 different genes were identified and classified into five categories according to the putative gene function of the highest-quality BLASTp alignment (Table 2).

GAM strains affected in metabolic-related genes (9 strains, 35.6% of the total) showed CI_{BM-RE} values ranging from approximately $10^{-3}$ to $10^{-1}$ (Fig. 1). Four of these mutants showed disruptions in genes related to the biosynthetic pathways of purine (GAM18, purB) and the amino acids glutamate (GAM9, gltB) and leucine (GAM12 and GAM21, leuC). Strains GAM9, GAM12 and GAM21 were unable to grow on minimal SSM plates (Table 2) but grew on reach LB medium (Fig. 1) and were therefore considered auxotrophs. Our screening also identified genes encoding enzymes involved in the $\beta$-oxidation of fatty acids (fadE and fadD), the degradation of the aromatic amino acids phenylalanine and tyrosine (hmgB), the catabolism of hydroxybenzoate (pcaC) and the assimilation of acetate, acyclic terpenes and leucine (aceA) (Table 2).

Transposon insertions in five of the 26 selected mutants were located in putative regulatory genes (CI_{BM-RE} values from $>10^4$ to approximately $10^1$). The interrupted genes in these strains are related to regulation of the response to changes in nitrogen balance (ntrB) (25), regulation of membrane functionality (colS) (26), metal resistance (copR) (27), and regulation of both alginate production and quorum sensing (algQ) (28,29). Higher CI_{BM-RE} values ($>10^{-2}$) were obtained for strains GAM6 (dppA) and GAM14 (kefA), which were affected in genes encoding components of putative transporters, i.e., a dipeptide ABC transporter and a mechanosensitive channel involved in potassium efflux, respectively. Other genes interrupted by the transposon in GAM strains encode hypothetical proteins, nucleic acid-related proteins, such as RecB and
DbpA, a putative dynamin-like protein (DLP), a GGDEF/EAL domain-containing protein and a peptidase M23 family protein (Table 2).

DNA Context Analysis of Transposon Insertions in the Genome of *Pseudomonas pseudoalcaligenes* AVO110. The genetic context surrounding each of the sequences interrupted by the transposon in GAM strains was analyzed. Special attention was given to the possible polar effect of the transposon insertions in the transcription of downstream genes (30). Out of the 24 open reading frames (ORFs) interrupted by the transposons, nine were considered to possibly form operons (Table 2).

Our STM approach identified several genes previously highlighted for their relevant role in rhizosphere colonization and competitiveness, such as *recB* (GAM22), *colS* (GAM24) and *algQ* (GAM26) (31,32). Taking into account that the biocontrol ability of *P. pseudoalcaligenes* AVO110 has been related to its ability to efficiently colonize avocado roots and *R. necatrix* hyphae, these three mutants were selected for further genetic characterization. Strains GAM2 and GAM3, which contain the transposon in genes encoding a putative DLP and a GGDEF/EAL domain-containing protein, respectively, were also selected for these analyses. These two genes especially caught our attention due to their possible role in bacterial-host interactions. Bacterial DLPs have been suggested to play a role in membrane remodeling under environmental stresses (33,34). On the other hand, GGDEF/EAL domain-containing proteins have been previously related to rhizosphere colonization in *Pseudomonas putida* (35) and *P. fluorescens* (36).

**Fig. 2** summarizes the genetic context of the genes interrupted by the transposon in the five selected GAM strains. RAST annotation identified the protein encoded by the gene interrupted in GAM2 as an ortholog of *E. coli* LeoA, a GTPase domain-containing
DLP (37). This DLP-encoding gene was found to overlap one bp with its upstream gene, encoding a hypothetical protein. In accordance with the localization of the leoABC operon in an E. coli H10407 pathogenicity island (38), a gene encoding a phage integrase was detected downstream this putative operon in AVO110 (Fig. 2).

The gene interrupted by the transposon in GAM3 possibly forms an operon with four ORFs showing homology (coverage 95%-100%; identity 59%-77%) with several Pseudomonas spp. genes involved in flagellar motility/chemotaxis (cheR, cheW, cheA and cheB). On the other hand, and in agreement with the operon organization of recB, recC and recD in other bacteria (39), the AVO110 recB gene, interrupted in GAM22, was also found in the proximity of recC and recD. This was also the case for the colS gene, interrupted in GAM24, which encodes the sensor protein of the two-component system ColR/ColS. Although insertion of the transposon in GAM26 possibly affects a single locus (algQ), other genes related to the production of exopolysaccharides (algZ, algR and algP) and porphyrins (hemC, hemD, hemX and hemY) were found in the proximity of this gene (Table 2, Fig. 2).

Modulation of Transcript Levels in wild-type Pseudomonas pseudoalcaligenes AVO110 After Transfer to Fungal Exudate-Containing Medium. Expression of the genes interrupted by the transposon in the five selected GAM strains (GAM2, GAM3, GAM22, GAM24 and GAM26) was analyzed after transfer of wild-type P. pseudoalcaligenes AVO110 cells to fungal exudate-containing medium (BM-RE). For this purpose, AVO110 cells were grown in LB medium to an OD 600 of 0.5, washed, and then transferred to BM-RE medium. Samples for RNA extraction were taken immediately after transfer to BM-RE (time 0) and after 4 h and 24 h of incubation. Fig. 3 shows the expression of these five genes, normalized to the housekeeping gene rpoD,
in BM-RE medium relative to their expression at time 0 (relative fold changes). While the levels of recB transcripts did not change after transfer of AVO110 cells to BM-RE medium, the transcript levels of the DLP-encoding gene were reduced by half at both 4 h and 24 h after transfer to BM-RE medium. However, the transcript levels of the remaining three genes increased after transfer to fungal exudate-containing medium. The sharpest increase in transcript levels was observed for the GGDEF/EAL domain-encoding gene, which showed values that were approximately 32 and 17 times higher at 4 h and 24 h after transfer to the BM-RE medium, respectively (Fig. 3).

Altered Colonization of R. necatrix Mycelia by Pseudomonas pseudoalcaligenes GAM Strains. The competitiveness of the selected P. pseudoalcaligenes GAM strains after inoculation over R. necatrix mycelia was tested in competition assays with the wild-type strain. GAM strains (Nf\(^R\) Km\(^R\)) could be differentiated from P. pseudoalcaligenes AVO110 (Nf\(^R\)) using plates containing kanamycin (Km). Mixed inocula of AVO110 and each of the five GAM mutants were prepared and used to inoculate R. necatrix mycelia grown on BM plates. Bacteria were recovered from the fungal hyphae 6 days after incubation at 25°C. After this period, the total number of P. pseudoalcaligenes cells recovered from the fungal mycelia ranged in all cases between \(10^4\)-\(10^5\) CFU per g of mycelia. Considering that P. pseudoalcaligenes AVO110 is unable to grow in this medium in the absence of R. necatrix mycelia or fungal exudates (1), the number of CFU recovered for each of the strains reflects their growth and survival under the influence of the fungal mycelia. While strain GAM3 (GGDEF/EAL mutant) was more competitive than AVO110 (CI > 1) during colonization of fungal hyphae, strains GAM2 and GAM24, which were affected in the DLP-encoding gene and the colS gene, respectively, were out-competed by the wild-type strain (CI >1).
Finally, strains GAM22 (recB) and GAM26 (algQ) showed CI values that were not significantly different from one, indicating that they were as competitive as AVO110 (Fig. 4).

**Altered Colonization of Avocado Roots by *Pseudomonas pseudoalcaligenes* GAM Strains.** Wild-type *P. pseudoalcaligenes* AVO110 was tagged with a mini-Tn7 derivative (mini-Tn7Km) encoding a Km resistance gene and the *gfp* gene (AVO110-Km, Table 1). *P. pseudoalcaligenes* GAM strains, which were already resistant to Km due to the insertion of the mini-Tn5Km2 derivative, were tagged with mini-Tn7Gm, which encodes a gentamicin (Gm) resistance gene and the *gfp* gene (Table 1). Competition assays between AVO110-Km and each of the Gm-tagged GAM mutants (GAM-Gm) during growth on LB medium revealed that all GAM-Gm strains were as competitive as AVO110-Km, suggesting that expression of double antibiotic resistance by GAM-Gm strains does not affect bacterial fitness under these conditions (Fig. S2).

*P. pseudoalcaligenes* AVO110-Km and each of the constructed GAM-Gm strains (Table 1) were individually inoculated in the roots of commercial 6-month-old avocado seedlings (cv. Walter Hole). At 7, 15, 30, 48 and 72 days after inoculation, bacteria were extracted from the avocado roots and plated in LB-Km (AVO110-Km) or LB-Km-Gm (GAM-Gm strains). Codification of the *gfp* gene within the miniTn7 derivatives used to construct these strains facilitated tracking of *P. pseudoalcaligenes* colonies and their differentiation from other rhizosphere bacteria. All strains were able to establish in the root system of avocado seedlings during the first week postinoculation, reaching approximately $10^7$ CFU g$^{-1}$ fresh root at 7 days postinoculation (dpi). Thereafter, bacterial counts for AVO110-Km slowly decreased to about $10^6$ CFU g$^{-1}$ fresh root at the end of the experiment (72 dpi). These results are in agreement with the previously
reported persistence of wild-type AVO110 on the roots of avocado plants (12), indicating that root colonization by AVO110-Km is not affected by the mini-Tn7Km transposon. A similar ability to establish and survive in the avocado rhizosphere was observed for strain GAM2-Gm (DLP mutant). In contrast, a faster decline in CFU counts was observed for strains GAM22-Gm (recB mutant), GAM24-Gm (colS mutant) and GAM26-Gm (algQ mutant), which showed $10^4$-$10^5$ CFU g$^{-1}$ fresh root at 72 dpi. On the other hand, strain GAM3-Gm (GGDEF/EAL mutant) established in the avocado rhizosphere at slightly higher cell counts than AVO110-Km at almost all sampling times. However, both strains reached a similar cell density at the end of the experiment (Fig. 5). Together, these results revealed differences between the colonization ability of the wild-type derivative AVO110-Km and some of the analyzed GAM-Gm strains.

While strain GAM3-Gm (GGDEF-EAL mutant) showed a slightly higher ability than AVO110-Km to persist in the avocado rhizosphere, GAM strains affected in recB, colS and algQ showed a lower colonization ability than the wild-type derivative.

The GGDEF/EAL Domain-Encoding Gene Interrupted in GAM3 Forms a Transcriptional Unit with a cheRWAB Gene Cluster. The increased fitness of the GAM3 mutant during colonization of R. necatrix hyphae (Fig. 4) and avocado roots (Fig. 5), together with the localization of the gene interrupted in this strain (GGDEF/EAL) in the proximity of a gene cluster (cheR, cheW, cheA and cheB) possibly involved in flagellar motility/chemotaxis (Fig. 2), prompted us to investigate whether all these genes are co-expressed as an operon that could be involved in the determination of these phenotypes. Co-transcription was analyzed by real-time polymerase chain reaction (RT-PCR) assays performed using RNA samples isolated from LB-grown P. pseudoalcaligenes AVO110 cells. Amplification of the intergenic
regions located between the sequential ORFs, with the exception of the regions upstream cheR and downstream the GGDEF/EAL domain-encoding gene (Fig. 2). revealed that all these genes are co-transcribed (Fig. 6).

The chemotactic response of wild-type P. pseudoalcaligenes AVO110 and the GAM3 mutant towards R. necatrix exudates was tested in minimal medium plates (0.2% agar). However, no chemotactic movement was observed for any of these strains under the conditions tested (data not shown). We also tested the swimming ability of the GAM3 mutant in comparison to the wild-type strain using 0.3% KB agar plates. No significant differences in the diameters of the swimming halos (measured after 24 h, 48 h and 72 h) were found between the strains (data not shown), indicating that motility of the GAM3 mutant was not affected under the conditions tested.

DISCUSSION

STM has been successfully applied to identify virulence genes in a vast number of human (40,41,42), animal (43,44) and plant bacterial pathogens (45,46,47,48). In addition, this strategy has been used for the identification of genes required during the interaction of beneficial bacteria with plants (49). Here, we report the application of STM to the interaction established between a biocontrol bacteria and a fungal phytopathogen. Although complete coverage of the P. pseudoalcaligenes AVO110 genome was not achieved in this study, STM libraries composed of a lower number of mutants have been successfully applied for the identification of virulence factors in bacterial plant pathogens (47,50) and animal pathogens (51).
Metabolic-Related Genes Required for Growth of AVO110 in *R. necatrix* Exudates. Nine out of the 26 (approximately 35%) GAM strains analyzed had interruptions in genes related to the metabolism of diverse compounds (Table 2). Strains GAM9 (*gltB*), GAM12 and GAM21 (both interrupted in *leuC*) contain disruptions of genes related to the biosynthetic pathways of the amino acids glutamate and leucine and were considered auxotrophs. These results, which are in agreement with the selection of auxotrophic strains in other STM studies (44,46,47), suggest that these two amino acids are limiting for growth of *P. pseudoalcaligenes* in *Rosellinia* exudate-containing medium. However, we cannot rule out that mutants with disruptions in the biosynthetic pathways of other amino acids also displayed limited growth in this medium, as they were either not represented in our library or were discarded. In relation to bacterial responses to the fungal exudate, the chemotactic response of *P. putida* and *P. tolaasii* to *Agaricus bisporus* mycelial exudate has been shown to be mainly dependent on amino acids, including leucine and glutamate (13).

The *purB* gene (GAM18), which encodes an adenylosuccinate lyase involved in the biosynthesis of purines (52), has been described as essential for rhizosphere colonization by *Pantoea agglomerans* (53) and for infection thread formation and nodule development in *Lotus japonicus* induced by *Mesorhizobium loti* (52). In relation to the relevance of the purine biosynthetic pathways in bacterial-fungal interactions and, in agreement with our results, the *P. putida purM* gene was shown to be involved in colonization of fungal mycelia by IVET (23). Future identification of compounds secreted to the medium by *R. necatrix* should shed light on the specific nutrients required for the association established between *P. pseudoalcaligenes* and this fungal pathogen.
Transporters and Transcriptional Regulators involved in Growth and Survival of AVO110 in *R. necatrix* Exudates. The *kefA* gene, which is interrupted by the transposon in GAM14, encodes a potassium-efflux system involved in bacterial protection against the detrimental effects of electrophilic compounds via acidification of the cytoplasm (54). On the other hand, the interrupted gene in strain GAM6 (*dppA*), encodes a periplasmic dipeptide-binding protein required for dipeptide transport and chemotaxis (55). Despite the relevance of these genes in bacterial physiology, their role in bacterial interactions with fungi remains to be elucidated.

Several transcriptional regulators were found to be required for the growth and persistence of *P. pseudoalcaligenes* AVO110 in fungal exudate (Table 2). Other transcriptional regulators have been related to bacterial adaptation and tolerance to adverse conditions generated in the proximities of fungal hyphae (14). For example, the *copRS* operon (GAM20) and the *ntrB* gene (GAM 8) have been involved in other bacteria with metal resistance and responses to changes in nitrogen balance, respectively (56,57). The possible role of the remaining transcriptional regulators identified in this study (*algQ* and *colS*) in the ability of AVO110 to survive in fungal exudates is discussed below.

AVO110 Genes Involved in Colonization *R. necatrix* Hyphae. The gene interrupted by the transposon in strain GAM2 (DLP-encoding gene) showed reduced transcript levels after exposing wild-type *P. pseudoalcaligenes* AVO110 to fungal exudates (Fig. 3). On the other hand, inactivation of this gene in GAM2 resulted in a reduced ability of this mutant to grow and survive in fungal exudates (Fig. 1) and to persist on *R. necatrix* hyphae (Fig. 4), indicating that this DLP is involved in the ability of AVO110 to establish a close association with *R. necatrix*.
The gene interrupted by the transposon in GAM3, which encodes a GGDEF-EAL domain-containing protein, is possibly involved in the metabolism of cyclic di-GMP (c-di-GMP). This second messenger has been reported to regulate a wide range of functions, including the switch between the planktonic and sessile lifestyles, bacterial adhesion and motility, responses to root exudate, colonization of host tissues and virulence (58,59,60,61). Moreover, we have shown that this gene is co-expressed with a cheRWAB cluster (Fig. 6), possibly involved in flagella-mediated chemotactic responses and motility (62). However, no differences in swimming motility were found between the GAM3 mutant and the wild-type strain. Inactivation of Rup4959, a P. putida GGDEF/EAL-domain containing protein induced by root exudate, did neither alter the swimming ability of the strain or its ability to interact with plants (35). On the other hand, and although expression of the cheRWAB genes is probably not affected by the transposon insertion in the GAM3 mutant (Fig. 2), inactivation of che-related genes generally results in alteration of flagella-mediated motility towards chemoattractants (62). Although we tested the chemotactic movement of wild-type P. pseudoalcaligenes AVO110 and the GAM3 mutant towards R. necatrix exudates, this response was not observed for either of the two strains under the conditions tested (data not shown).

Identification of rhizosphere and fungal exudate attractants involved in swimming chemotaxis and c-di-GMP signaling is needed to gain greater insight into the role of these mechanisms in biological control of fungal pathogens.

The gene interrupted by the transposon in strain GAM24 (colS) showed increased transcript levels after exposing P. pseudoalcaligenes AVO110 to fungal exudates (Fig. 3). In addition, this mutant showed a reduced ability to colonize both R. necatrix hyphae (Fig. 4) and the avocado rhizosphere (Fig. 5). The colS gene encodes a sensor element of the two-component system ColR/ColS, which is involved in bacterial outer
membrane permeability (31,63). In agreement with the reduced ability of strain GAM24 to colonize the avocado rhizosphere, a P. fluorescens colS mutant has been shown to be defective in competitive root colonization (31).

AVO110 Genes Required for Efficient Colonization of Avocado Roots but not for Establishment in R. necatrix Hyphae. Although strains GAM22 (recB mutant) and GAM26 (algQ mutant) were out-competed by the parental strain in BM-RE medium (Fig. 1), both mutants were as competitive as the wild-type strain during colonization of R. necatrix hyphae (Fig. 4), suggesting that these two genes are not essential during the physical interaction of P. pseudoalcaligenes AVO110 with fungal hyphae. However, their ability to colonize the avocado rhizosphere was reduced in comparison with the parental strain (Fig. 5). The recB gene encodes the helicase protein conforming the RecBCD holoenzyme, which is involved in homologous recombination and in repair of bacterial DNA damage. This enzyme has been related to efficient biofilm formation and host colonization (64,65). On the other hand, AlgQ regulates the production of alginate, a polysaccharide known to be involved in biofilm protection and colonization of plant tissues (28,66). In addition, AlgQ has been described as a global regulator in P. aeruginosa, upregulating siderophore synthesis and downregulating quorum sensing regulation (29).

In summary, this application of STM allowed us to identify genes of the bacterial biocontrol agent P. pseudoalcaligenes AVO110 that are required for growth and survival in the presence of fungal exudates, some of which are also essential for the colonization of fungal hyphae and/or plant root surfaces. Several metabolic pathways were highlighted as essential for the interaction of this bacterium with R. necatrix, such as those related to the metabolism of hydrocarbons, including acyclic terpenes, amino
acids, fatty acids and aromatic compounds. In addition, the relevance of a dipeptide transporter, metal resistance, protection against acidification of the cytoplasm and maintenance of nitrogen balance were also noted in this study as essential for bacterial survival under the influence of fungal exudate. Bacterial genes identified here as required for colonization of both fungal hyphae and plant roots are likely involved in membrane dynamics or the crosstalk between c-di-GMP signaling and chemotaxis. Finally, an additional set of two genes, which are perhaps related to responses to chemical stress and biofilm formation, were also identified as required for colonization of the avocado rhizosphere. Further functional characterization of these genes and of the compounds secreted by *R. necatrix* to the medium may promote a better understanding of the multitrophic interactions established among bacterial biocontrol agents, the plant rhizosphere and the mycelia of soilborne pathogens.

**MATERIALS AND METHODS**

**Bacterial Strains, Plasmids, Media, and Growth Conditions.** A pool of DNA sequence-tagged pUTmini-Tn5Km plasmids, *E. coli* strains CC118 λpir, S17-1 λpir, and DH5α and protocols for STM were kindly provided by D.W. Holden (Imperial College, London). The original bacterial strains and plasmids used in this study are listed in Table 1. *P. pseudoalcaligenes* strains were grown at 28°C in King’s medium B (KB) (67), lysogeny Broth (LB) medium (68), standard succinate medium (SSM) (69), or Super Optimal Broth (SOB) medium (70). *E. coli* strains were grown at 37°C in LB or SOB media. Solid and liquid media were supplemented, when required, with the following antibiotics (µg ml⁻¹) for *Pseudomonas/E. coli* strains: ampicillin (Amp): 100, Km: 25, nitrofurantoin (Nf): 50, Gm: 25 and cycloheximide (Ch): 50. The *P.*
*pseudoalcaligenes* derivatives selected in this study that contained a miniTn5Km2 transposon are listed in Table 2. The primers used in this study are listed in Table S1.

*R. necatrix* Rn400 was grown at 25°C on KB and stored at 4°C in water. Fungal mycelium was routinely replicated on KB plates to test viability every 6 months. BM minimal medium (4) was used for growing *R. necatrix* mycelia in order to obtain fungal exudate-containing medium (1) as described below.

**Preparation of BM Media Containing Fungal Exudates (BM-RE).** *R. necatrix* Rn400 was grown at 25°C on BM agar plates until the surface was completely covered by the fungus. To obtain liquid medium containing fungal exudates, fungal mycelium was collected from one plate, inoculated on 200 ml of minimal BM medium and incubated at 25°C for 2 weeks without shaking. Fungal mycelium was removed from the culture medium through filtration using sterile filter paper ALBET® in reams of 73 gr m² RM 2354252 (1). The final pH of the fungal exudate-containing medium (BM-RE) was 7.11.

**Generation of a Unique-Tag Marked Library of *Pseudomonas pseudoalcaligenes* Mutants.** A library of signature-tagged transposon mutants of *P. pseudoalcaligenes* was constructed as described by (71) with minor modifications. The pool of tagged pUTminiTn5Km2 vectors was transferred from *E. coli* S17-1 λpir to *P. pseudoalcaligenes* by plate conjugation mating as previously described (1). The transposition frequency of mini-Tn5Km2, which confers resistance to Km, was 6.8 x 10^6 transconjugants/receptor in the genome of *P. pseudoalcaligenes* AVO110 using biparental mating delivery. The constructed random transposition library consisted of 38 different 96-well microtiter trays, containing a total of a total of 3,408 *P. 
Pseudoalcaligenes mutants. Individual colonies were challenged on LB-Amp plates to discard P. pseudoalcaligenes transconjugants harboring the plasmid vector.

Agarose gel electrophoresis and other standard recombinant DNA techniques were performed as described previously (72). Genomic DNA was extracted using the Jet Flex Extraction Kit (Genomed, Löhne, Germany) according to the manufacturer’s instructions. Single and random transposon insertions into the genomes of the mutant strains were confirmed by Southern hybridization (73).

Colony Blots. To fix total DNA from the colonies, overnight cultures of P. pseudoalcaligenes mutant strains grown on LB-Km microtiter plates were transferred onto nylon membranes placed on LB-Km agar plates using a 48-pin replicator (Sigma-Aldrich, Inc., St. Louis, MO, USA). Colony blots were performed as previously described by (17).

STM Screening. STM screening was carried out by testing 76 pools of ≤45 mutants mixed with a negative control (wild-type P. pseudoalcaligenes AVO110) and a positive control (a miniTn5Km2-tagged AVO110 derivative selected by its ability to growth and survive on both LB and BM-RE). The input pools were generated by mixing 100 µl of cultures grown for 24 h at 28°C on microtiter plates containing LB (two wells per pool incubated with the negative control) and LB-Km (AVO110 mutants and one well per pool incubated with the positive control). Next, the mixtures were washed twice with NaCl 0.9% and adjusted to an OD₆₀₀ of 0.1 (approximately 10⁶ CFU ml⁻¹). Afterwards, 100 µl of these suspensions was used to inoculate 5 ml of BM medium amended with 5 ml of BM-RE (10³-10⁴ CFU ml⁻¹), which was incubated at 28°C and 225 rpm. After two days, mutant cells were recovered from the media (10⁶-10⁷ CFU ml⁻¹) to generate the
output pool (17). Finally, the 40-bp probes were purified using MiroSpin™ G-50 columns (GE Healthcare, Buckinghamshire, UK). DNA hybridizations on colony blots to 32P-labeled probes were carried out as described by (17). A schematic representation of the STM selection process is shown in Fig. S1.

Determination of Transposon Insertion Sites. Genomic DNA from selected mutants was digested with EcoRI and ligated into pBluescript II SK digested with the same restriction enzyme. Ligation reactions were used to transform DH5α by heat shock (70), and single Km-resistant colonies were selected. Plasmids showing DNA fragments of at least 1.8 Kb were purified using a NucleoSpin Plasmid Quick Pure kit, and the DNA regions flanking the transposons were sequenced using primer P7 (Table S1) (71). Automated DNA sequencing was performed by SECUGEN (Madrid, Spain). The raw sequences were analyzed by general BLASTn searches against NCBI-deposited sequences.

The DNA context surrounding each of the genes interrupted by the transposons in GAM strains was analyzed using the Artemis 13.2.0 genome browser/editor (74) and BLAST searches (http://blast.ncbi.nlm.nih.gov/Blast.cgi). The sequences surrounding the transposon in a selection of GAM strains were deposited in National Center for Biotechnology information (NCBI) (http://www.ncbi.nlm.nih.gov/) under the following accession numbers: GAM2 (KX863700), GAM3 (KX858709), GAM22 (KX906975), GAM24 (KX858711), and GAM26 (KX858710).

Draft Genome Sequencing and Annotation. Genomic DNA from P. pseudoalcaligenes AVO110 was extracted from bacterial cells grown overnight in LB medium supplemented with Nf at 25°C. DNA was extracted using a Genomic DNA
Purification JETFLEX kit (Genomed GmbH, Löhne, Germany) following the manufacturer’s instructions. The DNA sample was further purified by a first extraction with phenol:chloroform:isoamyl alcohol (25:24:1), followed by a second extraction with chloroform: isoamyl alcohol (24:1). DNA was precipitated with 1/10 volume of 3 M sodium acetate and two volumes of 100% ethanol and resuspended in MilliQ water. NanoDrop measurements gave a concentration of 315 ng µl\(^{-1}\) (315 µg of DNA in total) with an A260/A280 of 1.79. The draft genome of *P. pseudoalcaligenes* AVO110 was sequenced using the Illumina Hiseq 2000 platform at BGI Tech Solutions Co., Ltd. (Hong Kong), and pair-ended reads with insert sizes of 500 bp were assembled using SOAP de novo software (75,76). Statistics regarding the assembly results are summarized in Table S2. Automatic annotation of the draft genome was obtained using the RAST server (77).

In vitro Competition Assays. *In vitro* competition assays were performed as previously described (78). Competitive indexes (CIs) were calculated by dividing the output ratio (CFU mutant:CFU wild-type) by the input ratio (CFU mutant:CFU wild-type). The LB and BM-RE competition indexes shown (CI\(_{LB}\) and CI\(_{BM-RE}\), respectively) are the mean of three independent experiments ± standard deviation. Data were analyzed as described below (see Statistical Analysis).

Mini-Tn7 Tagging of GAM Strains. The *gfp* delivery plasmid pBK-miniTn7-Gm-*gfp* and the helper plasmid pUX-BF-13 were introduced into GAM strains by tetraparental mating as described by (79). Cells of the corresponding recipient strain, *E. coli* donor strain XL1-Blue (pBK-miniTn7(Gm)\(_{PA104D03-}gfp\)), helper strains *E. coli* SM10 (pUX-BF13) and *E. coli* HB101 (pRK600) (Table 1) were mixed at a 3:1:1:1 ratio,
respectively. PCR analyses on chromosomal DNA of the corresponding *gfp*-tagged strains (Table 1) were performed to determine whether miniTn7 insertion in GAM strains (Table 2) occurred at a specific site without gene disruption. In agreement with data reported for other *Pseudomonas* strains, PCR products that were approximately 150 bp long were amplified from derivative GAM strains using primers Tn7-*glmS* and Tn7-R109 (80, 81).

**RT-PCR and quantitative RT-PCR Assays.** For RT-PCR, *P. pseudoalcaligenes* AVO110 cells were grown in LB to an optical density at 600 nm (OD$_{600}$) of 0.5. The cells were pelleted and stored at -80°C. Total RNA was extracted using the TriPure isolation reagent (Roche Applied Science), as described previously (82). RNA concentration was determined spectrophotometrically, and its integrity was assessed by agarose gel electrophoresis. Total RNA was treated with a TURBO DNA-free kit (Applied Biosystems, Foster City, CA, U.S.A.), as detailed by the manufacturer’s instructions. DNA-free RNA was reverse transcribed using random hexamers included in the iScript cDNA synthesis kit (BioRad, Hercules, CA, U.S.A.). cDNA was used as a template to amplify intergenic regions by PCR, using GoTaq polymerase (Promega) and the primers indicated in Table S1. PCR products were analyzed by 1% agarose gel electrophoresis.

Quantitative real-time PCR (qRT-PCR) assays of *P. pseudoalcaligenes* AVO110 genes were performed using iQ™ SYBR Green Supermix (BioRad; California, USA) as follows. *P. pseudoalcaligenes* AVO110 was grown overnight in LB. The next day, the cells were diluted in 100 ml of LB medium to an OD$_{600}$ of 0.05 and grown to an OD$_{600}$ of 0.5. Cells were washed with NaCl 0.9% three times and inoculated in 100 ml of BM-RE to start the induction in this media. Three samples of 25 ml each were extracted
from this volume, and RNA extraction was carried out at time 0 and after 4 h and 24 h of incubation in BM-RE. RNA extraction and cDNA synthesis was performed as described above. Target cDNAs from the experimental samples were amplified in triplicate in separate PCR reactions using 0.3 M of each primer, GAM2QFwd/GAM2QRev for GAM2, GAM3QFwd/GAMQRev for GAM3, GAM4QFwd/GAM4QRev for GAM4, GAM22QFwd/GAM22QRev for GAM22, GAM24QFwd/GAM24QRev for GAM24 and GAM26QFwd/GAM26QRev for GAM26 (Table S1). The PCR amplicons were between 100 bp and 200 bp in length. Transcriptional data were normalized to the housekeeping gene rpoD from *P. alcaligenes* AVO110. After the normalization, expression fold changes at 4 h and 24 h were calculated with respect to the gene expression obtained right after the transfer to BM-RE medium (time = 0). qRT-PCR values are the mean of three biological replicates with three technical replicates ± standard deviation. Data were analyzed as described below (see Statistical analysis).

**Bacterial Colonization on Fungal Mycelia.** Colonization of fungal mycelia by GAM strains was determined in competition with the wild-type strain *P. pseudoalcaligenes* AVO110 as previously described by (22) with slight modifications (1). Agar discs obtained from PDA agar plates containing actively growing *R. necatrix* mycelia were placed onto BM agar plates, on which AVO110 (1) and GAM strains are not able to grow, covered with a cellophane layer and incubated for 10 days at 25°C. After this period, 1.5 ml of bacterial suspension previously washed twice with NaCl 0.9% and containing 10⁴-10⁵ CFU ml⁻¹ were equally distributed along the surface of each plate, and the plates were left to dry under a flow chamber for 2 h. After 6 days of incubation at 25°C, the cellophane layer containing actively growing fungal and bacterial strains
was placed into sterile plastic bags, weighed, transferred to a lab blender and homogenized for 2 min with 2 ml of sterile NaCl 0.9% to release the bacteria. Suspensions were serially diluted and plated on KB supplemented with Nf, Ch and Km (GAM strains) and Ch and Nf for total bacterial counts. Bacterial counts were obtained after 48-72 h at 25°C. *P. pseudoalcaligenes* wild-type strain counts were calculated as a ratio of bacterial total counts to GAM counts.

CIs were calculated as described above. The CIs shown are the mean of three technical replicates from three independent experiments ± standard error. Data were analyzed as described below (see Statistical analysis).

**Bacterial Colonization on Avocado Roots.** *P. pseudoalcaligenes* AVO110-Km, GAM2-Gm, GAM3-Gm, GAM22-Gm, GAM24-Gm, GAM26-Gm derivative strains were used. Six-month-old avocado seedlings of the commercial rootstock cv. Walter Hole (Brokaw nursery, Spain) were disinfected and inoculated with bacteria as previously described by (12) using suspensions containing $10^3$-$10^4$ CFU ml$^{-1}$. Plants were placed in non-sterile vermiculite and grown in a growth chamber at 24°C, 70% relative humidity, and 16 h daylight. Bacterial recovery from the roots was performed as follows. Three seedlings were removed from the vermiculite and processed independently at 7, 15, 30, 48 and 72 days of plant growth. Roots were separated from the plant, placed into sterile plastic bags, weighed and subsequently transferred to a lab stomacher and homogenized for 4 min with 2 ml of sterile PBS 0.1 M pH 7.2 per g of fresh root material. Suspensions were serially diluted and plated on LB supplemented with Nf and Gm for counts of GAM strains and Nf and Km for counts of the wild-type strain. Bacterial counts with appropriate colony morphology and antibiotic resistance, were obtained after 48-72 h at 25°C. Data represent the average of at least three
independent plants per sampling point ± standard error.

Statistical Analysis
Data were analyzed using SPSS software v.22 (Inc., Chicago, IL, USA). Competition index values resulting from in vitro competition assays and qRT-PCR values were analyzed using a Student’s t-test and the following null hypothesis: mean index was not significantly different from 1.0 (using p-values of 0.05). Competition index values obtained from bacterial colonization on fungal mycelia were subjected to one-way ANOVA followed by Tukey’s HSD test with the correction of Bonferroni (p= 0.05).

AUTHORS AND CONTRIBUTORS
AV, FMC and CR conceived the study; CP, IPM, AV, FMC and CR designed the experiments; CP, JICG, IPM and AP performed the experiments; CP, JICG, IPM, AP FMC and CR analyzed the data and interpreted the results; CP, IPM, FMC and CR drafted the manuscript with contributions from AP and AV; CP, JICG, IPM and AP prepared the Figures; and all authors read and approved the final manuscript.

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Table 1. Strains and plasmids used in this study

| Strains/plasmids | Relevant characteristics | Reference or source |
|------------------|--------------------------|---------------------|
| **Bacteria**     |                          |                     |
| Rosellinia necatrix Rn400 | Wild-type strain | (87) |
| GAM2-Gm         | GAM2 (Km\(^3\)) tagged with mini-Tn7Gm, also encoding the gfp gene. | This study |
| GAM22-Gm        | GAM22 (Km\(^3\)) tagged with mini-Tn7Gm, also encoding the gfp gene. | This study |
| GAM24-Gm        | GAM24 (Km\(^3\)) tagged with mini-Tn7Gm, also encoding the gfp gene. | This study |
| GAM26-Gm        | GAM26 (Km\(^3\)) tagged with mini-Tn7Gm, also encoding the gfp gene. | This study |
| Escherichia coli | hsdR17, supE44, recA1, endA1, gyrA46, thi, relA1, lac F' [proAB\(^R\), lacI\(^R\), lacZ M15::Tn10(Tc\(^R\))]. | (83) |
| XL-Blue         | F\(^-\) lacI\(^{-}\) Thr\(^{-}\) Leu\(^{-}\) Met\(^{-}\) Spc\(^{-}\) Tn7\(^{-}\) Sm\(^{-}\) (800lacZAM15, Δ(lacZYA-argF)U169, deoR, recA1 endA1, hsdR17(k\(_{+}\) m\(_{+}\)), phoA, supE44, λ, thi-1, gyrA96, relA1. | (70) |
| S17::apir       | thi pro hsdR recA RP4-2 (Tc::Mu Km::Tn7λ.Tc\(^R\) Stu\(^R\) ]-pir lysogen. | (84) |
| CC118::pir      | Δ(ara-leu) araD ΔlacX74 gaiE gaiK phoA20 thi-1 rpsE rpoB argE(Am) recA1, λ-pir lysogen, (Rif\(^R\)). | (85) |
| HB101           | Sm\(^R\) recA thi pro leu hsdR M\(^R\). Used for replication of the helper plasmid for RK600. | (86) |
| SM10::pir       | Thi-1 thr leu tonA lac Y supE recA::RP4-2-Tc::Mu, Km\(^R\), Apir. Used for replication of pUX-BF13. | (79) |
| **Fungi**       |                          |                     |
| Pseudomonas pseudoalcaligenes AVO110 | Wild-type strain | (12) |
| AVO110-Km       | AVO110 tagged with mini-Tn7-km, also encoding the gfp gene. | (1) |
| GAM2-Gm         | GAM2 (Km\(^3\)) tagged with mini-Tn7Gm, also encoding the gfp gene. | This study |
| GAM3-Gm         | GAM3 (Km\(^3\)) tagged with mini-Tn7Gm, also encoding the gfp gene. | This study |
| GAM22-Gm        | GAM22 (Km\(^3\)) tagged with mini-Tn7Gm, also encoding the gfp gene. | This study |
| GAM24-Gm        | GAM24 (Km\(^3\)) tagged with mini-Tn7Gm, also encoding the gfp gene. | This study |
| GAM26-Gm        | GAM26 (Km\(^3\)) tagged with mini-Tn7Gm, also encoding the gfp gene. | This study |

\(^a\)Strains GAM2, GAM3, GAM22, GAM24 and GAM26, which are *P. pseudoalcaligenes* AVO110 derivatives containing a mini-Tn5Km2 transposon encoding a kanamycin resistance (Km\(^3\)) gene, are described in Table 2.
Table 2. *Pseudomonas pseudoalcaligenes* AVO110 growth-attenuated mutants (GAM) selected by signature-tagged mutagenesis (STM).

| Name  | Gene Identity | Predicted Function                  | SSM<sup>b</sup> | ORF length | MiniTn5<sup>c</sup> insertion<sup>d</sup> | Possible polarity<sup>d</sup> |
|-------|---------------|-------------------------------------|------------------|------------|------------------------------------------|-----------------------------|
| **Metabolism** | | | | | | |
| GAM5  | hmgB           | Fumarylacetoacetase hydrolase       | +                | 783        | 292                                      | II                          |
| GAM9  | gltB           | Glutamate synthase large chain      | -                | 4,449      | 3,743                                    | I                           |
| GAM10 | aceA           | Isocitrate lyase                    | +                | 1,596      | 120                                      | III                         |
| GAM11 | fadE           | Acyl-CoA dehydrogenase              | +                | 2,448      | 840                                      | II                          |
| GAM12 | leuC           | 3-isopropylmalate dehydratase large subunit | -         | 1,428      | 62                                       | I                           |
| GAM17 | fadD           | Putative fatty acid CoA ligase      | +                | 1,611      | 324                                      | I                           |
| GAM18 | purB           | Adenylsuccinate lyase               | +                | 1,371      | 1,337                                    | II                          |
| GAM21 | leuC           | 3-isopropylmalate dehydratase large subunit | -         | 1,428      | 432                                      | I                           |
| GAM25 | pcaC           | 4-carboxymuconolactone decarboxylase| +                | 573        | 144                                      | I                           |
| **Transcriptional Regulation** | | | | | | |
| GAM8  | ntrB           | Histidine kinase, two-component sensor protein | +            | 1,077      | 173                                      | I                           |
| GAM15 | colS           | Histidine kinase, two-component sensor protein | +            | 1,281      | 1,240                                    | I                           |
| GAM20 | copR           | Transcriptional activator, two-component response regulator | +            | 675        | 292                                      | I                           |
| GAM24 | colS           | Histidine kinase, two-component sensor protein | +            | 1,281      | 1,001                                    | I                           |
| GAM26 | algQ           | Transcriptional regulator, anti-sigma factor | +            | 474        | 400                                      | III                         |
| **Transporters** | | | | | | |
| GAM6  | dppA           | Dipeptide ABC transporter           | +                | 1,842      | 4                                        | II                          |
| GAM14 | kefA           | Mechanosensitive channel, potassium-efflux system | +            | 3,369      | 987                                      | I                           |
| **Hypothetical proteins** | | | | | | |
| GAM1  | Hypothetical protein | | +  | 801 | 603 | III |
| GAM7  | Hypothetical protein | | +  | 354 | 116 | II  |
| **Other functions** | | | | | | |
| GAM2  | Dynamin-like protein, GTPase | | +  | 1,704 | 655 | I  |
| GAM3  | GGDEF-EAL domain protein | | +  | 3,570 | 1,741 | II |
| GAM4  | Peptidase M-23 | | +  | 855 | 73 | II |
| GAM13 | Intergenic region | | +  | - | - | nd |
| GAM16 | recB          | Exodeoxyribonuclease V, sub. B | +  | 3,543 | 3,520 | II |
| GAM19 | recB          | Exodeoxyribonuclease V, sub. B | +  | 3,543 | 3,515 | II |
| GAM22 | dbpA          | ATP-independent RNA helicase (ribosome assembly) | +  | 1,335 | 390 | I |

<sup>a</sup>Name of the strain and functional group to which it belongs. Light gray background indicates GAM strains selected for further characterization according to the predicted function of the gene interrupted by the transposon.

<sup>b</sup>Ability (+) or inability (-) to grow in standard succinate medium (SSM) minimal medium.

<sup>c</sup>MiniTn5<sup>k</sup> insertion point. Numbers indicate the exact position (bp) of the transposon in the disrupted gene, assuming the first nucleotide of the start codon = 1.
Possible polarity of miniTn5Km2 insertions in fitness-attenuated mutant (GAM) strains. Operon predictions were performed following the criteria defined by (30). Intergenic distances of < 40 nt between the genes interrupted by the transposons and the next downstream genes were considered to affect operonic pairs (I); distances > 40 but < 200 nt were considered to affect operonic pairs with intermediate probability (II); at distances > 200 nt, mutations were considered to affect probable single loci. Genes interrupted by the transposon located upstream of a gene transcribed in the opposite orientation were also included in this category (III). nd, not determined, the transposon was located within an intergenic region.
FIGURE LEGENDS

Fig. 1. Competition assays of *Pseudomonas pseudoalcaligenes* growth-attenuated mutants (GAM) strains. Competitive index values (CI) are shown for mixed inoculations of *P. pseudolacaligenes* and its derivative GAM strain. CI_{LB} (black bars), competition index values in lysogenic broth (LB) medium. CI_{BM-RE} (gray bars), competition index values in minimal BM medium supplemented with *R. necatrix* exudates (BM-RE medium). CI assays of GAM strains disrupted in metabolic-related (A) and non-metabolic-related (B) genes. CIs are the mean of three samples, and the error bars represent the standard deviation from the average. In all cases, CI_{BM-RE} is significantly <1.0 and significantly lower than CI_{LB}. Statistical analyses were performed using Student’s *t*-test (p = 0.05).

Fig. 2. DNA context of transposon insertions in *Pseudomonas pseudoalcaligenes* AVO110 growth-attenuated mutants (GAMs). Arrows indicate the direction of transcription and relative sizes of the genes in the genome of *P. pseudoalcaligenes* AVO110. Numbers within or over the arrows designate GenBank accession numbers. Gray arrows indicate the gene interrupted by the miniTn5 transposon. The inverted black triangles indicate the position where the transposon was integrated in the corresponding GAM mutant. Genes whose closest homologs are currently unnamed are indicated by the possible function of their corresponding encoded proteins as follows: DLP, dynamin-like protein; HP, hypothetical protein; GGDEF/EAL, GGDEF/EAL-domains protein; MP, membrane protein, and FKBP-I, FKBP-type peptidyl-prolyl cis-trans isomerase.

Fig. 3. Expression of five selected genes in wild-type *Pseudomonas pseudoalcaligenes* AVO110 after transfer to *Rosellinia* exudate-containing medium (BM-RE medium). The expression of the indicated genes (DLP, dynamin-like protein gene, GGDEF/EAL;
GGDEF/EAL domain-encoding gene; *recB, colS* and *algQ*) was measured by qRT-PCR in AVO110 at 4 h (black bars) and 24 h (white bars) after transfer to BM-RE medium. The fold change was calculated after normalization using the housekeeping *rpoD* gene as an internal control. After the normalization, expression fold changes at 4 h and 24 h were calculated with respect to gene expression obtained before the transfer to BM-RE medium (time = 0). qRT-PCR values are the mean of three biological replicates with three technical replicates. Bars represent the standard deviation from the average. Statistical analyses were performed using Student’s *t*-test (*p* = 0.05). Asterisks indicate values that deviate significantly from unity.

**Fig. 4.** Competition assays between *Pseudomonas pseudoalcaligenes* AVO110 growth-attenuated mutants (GAM strains) and the wild-type strain during colonization of *R. necatrix* mycelia. Competitive index (CI) values are shown for mixed inoculations of *P. pseudoalcaligenes* AVO110 and its derivative GAM strain in minimal medium BM plates covered by *R. necatrix* mycelia. The CIs shown are the mean of three samples, and the error bars represent the standard error. Bar values with different letters denote significant differences (*p* = 0.05) using one-way ANOVA followed by Tukey’s HSD test with the correction of Bonferroni.

**Fig. 5.** Colonization of avocado roots by *Pseudomonas pseudoalcaligenes* growth-attenuated mutants (GAM). Avocado seedlings (cv. Walter Hole) were inoculated with bacterial suspensions (10^3-10^4 CFU ml-1) of *P. pseudoalcaligenes* AVO110-kanamycin (Km) or its GAM-gentamicin (Gm) derivative strains (*Table 1*). Bacteria were recovered from the roots at 7, 15, 30, 48 and 72 days after inoculation and plated on LB supplemented with nitrofurantoin (Nf) and Gm for counts of GAM-Gm strains (closed circles) and Nf and Km for counts of the wild-type derivative strain AVO110-Km (open circles). Data represent the average of at least three independent plants per sampling.
point ± standard error.

Fig 6. Transcriptional analysis of the cheRWAB-GGDEF/EAL operon in *Pseudomonas pseudoalcaligenes* AVO110. Schematic representation of the intergenic regions amplified by RT-PCR (A); numbers in brackets indicate the sizes of the expected amplification products (numbered 1 - 6) in base pairs (bp). Gel electrophoresis (1.0 % agarose) of RT-PCR amplicons obtained using cDNA synthesized from RNA samples (B). M, molecular weight DNA marker (DNA ladder, Life Technologies). Numbers indicate amplification products shown in (A). The primer pairs used are detailed in Table S1.
cheB  cheA  cheR  cheW  GGDEF/EAL  dnaX
1
(629 bp) ( 292 bp) (297 bp) ( 293 bp) (272 bp)
2 3 4 5 6
1 2 3 4 5 6 M
A
B
100
300
500
1000
(444 bp)
bp