Virulence Attenuation of Pectobacterium carotovorum Using N-Acyl-homoserine Lactone Degrading Bacteria Isolated from Potato Rhizosphere

Esmaeil Mahmoudi1*, Badraldin Ebrahim Sayed Tabatabaei2 and Vittorio Venturi3
1Department of Plant Protection, Khorasgan Branch, Islamic Azad University, Isfahan, Iran
2Department of Agricultural Biotechnology, Isfahan University of Technology, Isfahan, Iran
3Bacteriology Group, International Center for Genetic Engineering and Biotechnology, Trieste, Italy
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Several soil bacteria were found to degrade N-Acyl-homoserine lactones (NAHLs), thereby interfering with the bacterial quorum sensing system. In this research, fifteen strains of NAHL degrading rhizobacteria were isolated from potato rhizosphere. Based on phenotypic characteristics and 16S rDNA sequence analyses, the strains were identified as members of genera Bacillus, Streptomyces, Arthrobacter, Pseudomonas and Mesorhizobium. All tested isolates were capable to degrade both synthetic and natural NAHL produced by Pectobacterium carotovorum subsp. carotovorum (Pcc) strain EMPCC. In quorum quenching experiments selected isolates, especially Mesorhizobium sp., were markedly reduced the pathogenicity of Pcc strain EMPCC in potato tubers and totally suppressed tissue maceration on potato tubers. These led to consider the latter as a useful biocontrol agent against Pectobacterium spp.

Keywords: Mesorhizobium, NAHLs, Pectobacterium carotovorum, Quorum sensing

Many bacterial species employ complex communication systems that link cell density and gene expression to a broad range of biological functions (Fuqua et al., 2001; Miller and Bassler, 2001). Such cell-to-cell communication, termed quorum sensing (QS), depends on the production, diffusion, and recognition of small signal molecules. Bacteria recognize the changes in their population density by sensing the concentrations of signal molecules, which are accumulated as bacterial cells proliferate. N-Acyl-homoserine lactones (NAHLs), also known as autoinducers, are widely conserved signal molecules that are present in the quorum-sensing systems of many gram-negative bacteria. Bacteria release, detect, and respond to accumulation of these signal molecules resulting in synchronizing expression of diverse biological functions including bioluminescence, antibiotic synthesis, biofilm formation, swarming, plasmid conjugation, siderophore production and production of virulence determinants (Whitehead et al., 2001). Since pathogenicity or pathogenicity-associated functions are controlled by QS in some major plant and animal pathogens, it has been proposed that the QS system could be an appropriate novel target for biological control agents or drugs in order to attenuate and limit virulence (Faure and Dessaux, 2007). Anti-QS mechanisms have been employed by inhibiting the synthesis or perception of the NAHL signal, as well as its enzymatic degradation by wild-type bacteria and eukaryotes, and genetically modified organisms (Molina et al., 2003; Rasmussen and Givskov, 2006).

The NAHL-degrading enzymatic activities were initially discovered in a few bacterial species (Dong et al., 2000; Leadbetter and Greenberg, 2000). However, they have now been reported in Proteobacteria belonging to the Agrobacterium, Bosea, Comamonas, Delftia, Ochrobactrum, Pseudomonas,Ralstonia, Sphingopyxis and Variovorax genera (d’Angelo-Picard et al., 2005; Flagan et al., 2003; Huang et al., 2003; Jafra et al., 2006; Leadbetter and Greenberg, 2000; Lin et al., 2003; Park et al., 2003; Uroz et al., 2003). In addition degrading activities have been originally reported in Actinobacteria and Firmicutes, such as Arthrobacter, Bacillus and Streptomyces (Dong et al., 2000; Lee et al., 2002; Park et al., 2003; Uroz et al., 2003). These NAHL-degrading bacteria were recovered from different environments such as soil, rhizosphere and abiotic biofilm.

The aims of this study were: (i) to isolate and identify bacteria that degrade NAHL molecules; (ii) to evaluate their NAHL degradation pattern; and (iii) to evaluate the efficacy of microbial-based NAHL degradation for biological control of the pathogen P. carotovorum subsp. carotovorum on potato plant.
Material and Methods

Bacterial strains, media and culture conditions. Bacterial strains were isolated from the rhizosphere of potato plants collected from different potato growing areas in Iran. *Pectobacterium carotovorum* subsp. *carotovorum* strain EMPCC (Laboratory of Plant Pathology, IAU- Science and Research Branch, Tehran, Iran) was used as a positive control for NAHL degradation agent in experiments. *Chromobacterium violaceum* CV026 (McClean et al., 1997) (provided by Vittorio Venturi, ICGEB, Area Science park, Italy) and *Agrobacterium tumefaciens* NT1 (Shaw et al., 1997) (provided by Yves Dessaux, CNRS, Gif-sur-Yvette, France) were used as the indicator strains for AHLS detection. The media used were Luria-Bertani (LB), King’s-B (KB) (Schaad et al., 2001) and AB minimal medium that was supplemented when required with 2% mannitol (Chilton et al., 1974) or with cycloheximide (50 μg/l). For growth of *E. coli* SM88, the media was supplemented with tetracycline (10 mg/l). All bacteria were grown at 27°C, except for biosensors and *E. coli* SM88, which were grown at 28°C and 37°C, respectively. All NAHLs standard used in this study were purchased from Sigma (Sigma-Aldrich, Inc., St. Louis, Mo., USA).

Isolation of Bacterial strains. Soil and potato root samples were collected from commercial potato fields located in Isfahan province, Iran. The original suspensions were made by adding 1 gram of either samples in 10 ml sterile distilled water. Serial dilutions are prepared in 10 ml of sterile 0.8% NaCl and a loopful of the most diluted suspensions were spread on King’s medium B and LB medium (1% tryptone, 0.5% yeast extract and 0.5% NaCl) containing 50 mg/mL cycloheximide. Plates were incubated in the dark at 27°C for 72h. The bacterial colonies with different morphologies were randomly picked from both media and the purified cultures were stored in glycerol medium at −80°C.

Screening of bacterial isolates for NAHL degradation activity. Screening for AHL degrading activities of potato bacterial isolates was performed as described by Morohoshi et al., 2009. Because NAHLs are sensitive to alkaline pH (Yates et al., 2002), all degradation assays were done in AB and LB media at pH 6.5. The N-hexanoyl homoserine lactone (C6-HSL) was used as first target molecule in this degradation assay. Bacterial strains were incubated into 5 ml LB medium containing 5mg/l C6-HSL on rotary shaker for 24 h at 27°C. Cells were removed by centrifuge at 2000 x g for 5 min. the culture supernatant (50 μl) was transferred into the wells of a 96-well plate. The full-grown culture of the CV026 biosensor was diluted 1:100 in fresh LB medium, and 500 μl of the diluted culture was inoculated into each well. A control experiment involving non-inoculated degradation medium processed as for the inoculated media was performed at the same time as the degradation assays. *E. coli* SM88 was used as positive NAHL degrading bacterium. After incubation for 24 h at 30°C on rotary shaker, the remaining C6-HSL was detected through violacein production by the *C. violaceum* reporter strain. To determine the NAHL degradation pattern of selected isolates, individual stock colonies were inoculated in 5 ml LB medium supplemented each with one of NAHLs standard with following concentrations: 6 mg/l (C8-HSL); 6.5 mg/l (C12-HSL) and 6.5 mg/l (C14-HSL). Cultures were incubated at 27°C for 24 h on rotary shaker. The control experiment with non-inoculated degradation medium was performed to assess degradation assays. Following this, bacterial cells were removed by centrifugation at 2,000 × g for 5 min and assay was performed as described above except for long chain NAHL, that NT1 strain was used as biosensor.

NAHL production by *Pc.* subsp. *carotovorum* EMPCC. *Pc.* subsp. *carotovorum* EMPCC was streaked as homogeneous line on LB medium and biosensor strain, *C. violaceum* CV026, was spotted at a distance of 6-7 mm from the EMPCC line. After incubation at 28°C for 24 h, appearance of violet pigment in CV026 colony revealed the production of violacein by CV026 as well as production of NAHL by EMPCC.

Degradation ability of natural NAHL produced by *Pc.* subsp. *carotovorum* EMPCC. *Pc.* subsp. *carotovorum* strain EMPCC was inoculated in 5 ml LB medium at 27°C by shaking for 24 h. The bacterial suspension was centrifuged at 2,000 × g for 5 min. Culture supernatant was extracted twice adding an equal volume of ethyl acetate. The residue was added with fresh LB medium and all test strains (Table 1) were inoculated in this combined medium. Bacterial cultures were incubated at 27°C for 20 h (for SM88 at 37°C) and NAHL residue was detected as described by Shaw et al. (1997).

Identification of the selected isolates. To confirm preliminary identification of bacterial isolates determined by some phenotypic characteristics (Schaad et al., 2001), the selective strains were subjected to PCR analysis. The DNA coding regions for the 16S rRNA of each isolate were amplified by PCR using the universal primers pA (5'-
AGAGTTTGATCCTGGCTCAG) and pH (5’-AGGAGGT-
GATCCAGCCGCA), which allowed the amplification of
almost the entire gene (Bruce et al., 1992). DNA extraction
for strains was performed as described previously by
Manzano et al. (2003). Polymerase chain reactions were
performed in a total reaction volume of 50 μl containing 1X
PCR buffer, 100 μM of each dNTP, 1.5 mM MgCl
2
, 0.1 μM primers, 100 ng of DNA extract and 1 U of Taq DNA
Polymerase (Cinagene, IRIB). The following temperature
cycle was used: an initial denaturation step of 5 min at 95
°C
followed by 30 cycles of 1 min denaturation at 95
°C, 1 min
annealing at 56
°C and 1 min 30s extension at 72
°C and a
final extension step of 5 min at 72
°C. The amplification
yielded a product of ca. 1500 bp, which was analyzed by
electrophoresis on 0.8% agarose gel and then by staining
with ethidium bromide (Sambrook et al., 2001). The result-
ing PCR products were sequenced by an BigDye Terminator
and ABI Prism 3700 Genetic Analyzer (Macrogen, World
Meridian Venture Center, Korea), and at least 400 bp were
subjected to the BLAST analysis within the NCBI database.
Some sequence comparisons authorized identification of
isolates at the species level and for others the only genus
level was retained for homogeneity.

**Ability of the tested strains to attenuate potato tissue
maceration by P. carotovorum subsp. carotovorum.** The
assay was performed on potato tubers (cv. Agria) as
described by Lojkowska et al. (1995). Potato tubers were
washed and surface sterilized by two consecutive incubations
with sodium hypochlorite (1% chlorine deg.), extensively
rinsed with sterile water. The tubers were dried under sterile
conditions and then were sprayed with 70% ethanol. They
were dried again and were kept for co-inoculation method.
Strains used in this assay were P. carotovorum
subsp. carotovorum
EMPCC (as pathogen) and AM51, S5 and
DMS133 as biocontrol agents (quenchers). Strains were
cultivated overnight at 27
°C in LB medium, suspended and
diluted in sterile 0.8% NaCl. Each tuber was inoculated
with 20 μl of bacterial suspension including pathogen,
pathogen with the quencher and quencher alone. Four
potato tubers were used for each combination of strains.
The experiments were repeated twice. After inoculation, the
potato tubers were incubated in a moisture chamber (over
90% humidity) at 25
°C. Three days after infection, the
tubers were cut in the middle and the results were assessed
by visual inspection and photographed.

**Results**

Isolation and Screening of degrading NAHLs bacteria.
Among forty-five rhizospheric soils and root samples,
eighty-four individual colonies with different morphologies

| Strains | rrs sequencing identification | GenBank Acc. no. | Gram | Colony and cell morphology | Degradation ability* |
|---------|-------------------------------|-----------------|------|----------------------------|----------------------|
| S6      | Bacillus sp.                  | EU977693.1      | +    | White, rod shape, motile   | ++                   |
| S22     | Bacillus sp.                  | HM748447.1      | +    | White, rod shape, motile   | ++                   |
| S27     | Bacillus sp.                  | HM776218.1      | +    | White, rod shape, motile   | ++                   |
| AM35    | Bacillus sp.                  | HM748447.1      | +    | White, rod shape, motile   | ++                   |
| AM38    | Bacillus sp.                  | EU240440.1      | +    | White, rod shape, motile   | ++                   |
| EA60    | Bacillus sp.                  | FJ66758.1       | +    | White, rod shape, motile   | ++                   |
| EA73    | Bacillus sp.                  | AY948211.1      | +    | White, rod shape, motile   | ++                   |
| EA85    | Bacillus sp.                  | D26185.1        | +    | White, rod shape, motile   | ++                   |
| DMS133  | Bacillus sp.                  | HM188452.1      | +    | White, rod shape, motile   | ++                   |
| S5      | Mesorhizobium                 | AF410896.1      | –    | White, rod shape, motile   | ++                   |
| EA113   | Streptomyces sp.              | HM748050.1      | +    | White, like filamentous, non-motile | ++                   |
| AM51    | Arthrobacter sp.              | AY444858.1      | +    | White to grayish, rod, non-motile | ++                   |
| AM43    | Arthrobacter sp.              | AY731366.1      | +    | White to grayish, rod, non-motile | ++                   |
| S15     | Arthrobacter sp.              | AY635865.1      | +    | White to grayish, rod, non-motile | ++                   |
| EA101   | Pseudomonas sp.               | AJ969084.1      | –    | White, rod, fluorescent on King-B, motile | ++                   |
| SM88    | E. coli                       | –               | –    | White, rod shape, motile   | ++                   |

a: BLAST homology searches (GenBank Acc. no.)
b: NAHL degrading ability was performed as described in methods.
c: P. car. subsp. carotovorum strain EMPCC used as NAHL production source, +: low degrading ability, ++: high degrading ability
were screened for NAHL degradation. Application of the pigment induction-based indicators, *Chromobacterium violaceum* CV026 and *Agrobacterium tumefaciens* NT1 for short chain and long chain carbon NAHLs respectively, allowed us to screen feasibly all NAHL degraders as described by McClean et al., 1997; Shaw et al., 1997. The results revealed that fifteen isolates completely degraded 5 mg/l of C6-HSL after 24 h and violacein induction was inhibited in the CV026 biosensor (Fig. 1). In order to characterize NAHL degradation patterns of the selected isolates, four synthesized unsubstituted NAHL and crude cell culture extract of *P. carotovorum* subsp. *carotovorum* strain EMPCC as natural NAHL were used. Results revealed that all tested strains degrade all NAHLs types under same conditions with high or low degradation activity (Table 1). In fact, the degradation properties of the various strains differed with respect to their substrate preferences. For instance, strains S6, S22, S27 and AM35 completely degraded all NAHLs as well as culture extract of EMPCC after 24 h (Fig. 2). These isolates were identified as *Bacillus* spp., which are known to have high NAHL degrading activity (Dong et al., 2000 and 2002). The other four strains namely AM51, AM43, S15 and EA101 could hardly degrade 5 mg/l C6 and C8-HSL after 24 h.

Therefore, reminiscent of C12 and C14-HSLs in cell culture supernatant and in turn, weak blue pigment production by NT1 biosensor, revealed their inability to degrade 6 mg/l of long chain NAHLs completely. Despite complete degradation of long chain NAHLs (C12 and C14-HSL) by *Streptomyces* sp. EA113, a trace of violacein production was induced when CV026 cultured on supernatant of C6 and C8-HSL medium. In addition, all tested strains excluding EA101 and EA113, completely degraded natural NAHL produced by *P. carotovorum* subsp. *carotovorum* EMPCC (Fig. 3, Table 1). The results described above indicated that all tested isolates were able to degrade various NAHLs with different abilities.

**Identification of the NAHL degrading isolates.** Bacterial

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**Fig. 1.** Detection of NAHL-degrading isolates. The NAHL-degrading isolates were detected by inhibition of synthesis of violacein by *C. violaceum* CV026 in the presence of C6-HSL at 6 mg/l. The fifteen NAHL-degrading isolates are numbered from 1 to 15. C (control) degradation assay performed without bacteria. The DH is *E. coli* DH5α (SM88) carrying aiiA lactonase encoding gene was used as positive NAHL degrading bacterium. Numbers 1 to 15 encoded as S5, S6, S15, S22, S27, AM35, AM38, AM43, AM51, EA60, EA73, EA85, EA101, EA113 and DMS133, respectively. The picture was taken after 24 h incubation.

**Fig. 2.** (A-1) NAHL production by *P. carotovorum* subsp. *carotovorum* strain EMPCC. Test strain was streaked as a homogeneous line on LB medium and biosensor strain, *C. violaceum* CV026, was spotted at 10 mm from the test strain. Development of violet pigment in CV026 colonies revealed the production of violacein by CV026 as well as production of NAHL by EMPCC. (A-2) *Pseudomonas fluorescens* strain PS1 was used as non-NAHL producing control. (B) Thin layer chromatogram for degradation of NAHL produced by *P. carotovorum* subsp. *carotovorum* EMPCC. Test strain was inoculated into LB medium and after 24 h incubation at 27°C, free cell culture supernatant was extracted with ethyl acetate. The extract was added into fresh LB medium and selected strains (DME133 and SM88) were cultured in it. After 24 h, culture supernatant was extracted with ethyl acetate and NAHL disappearance was revealed using the biosensor strain *Chromobacterium violaceum* CV026 as described in methods.
Biocontrol of P. carotovorum subsp. carotovorum by NAHL degrading bacteria. The biocontrol activity of tested isolates (Table 1) was performed against P. carotovorum subsp. carotovorum EMPCC, with this knowledge that its virulence is regulated by QS with 3-oxo-C6HSL as an essential signal (Dong et al., 2002). As shown in Fig. 3, the direct inoculation of P. carotovorum EMPCC caused extensive tissue maceration in 48 h incubation time. Whereas co-inoculation of EMPCC with NAHL degrading bacteria led to substantial reduction in tissue maceration compared to the pathogen alone (Fig. 3). In this regard, the inhibition activity of the B. subtilis strain DMS133 was more pronounced than other tested isolates. In contrast, the ability of EMPCC to macerate potato tubers tissue was attenuated by Arthrobacter sp. AM51 and significantly reduced by Mesorhizobium sp. S5 when co-inoculated with EMPCC at 10^6 cfu/ml (Fig. 3).

Discussion

The biological degradation of quorum sensing signal molecules is an important way in interrupting expression of regulated by quorum-sensing. In screening for NAHL degrading bacteria it was found that major soil and potato root surface bacteria which are taxonomically diverse are able to degrade NAHLs signal molecules. According to previous results (Dong et al., 2000; Angelo-Picard et al., 2004 and Jafra et al., 2006), 10−20% of total cultivable bacteria in soil are able to degrade NAHLs. This study allowed the isolation of several NAHL degrading strains from potato rhizosphere; 15/84 isolates were identified as Mesorhizobium, Pseudomonas, Bacillus, Streptomyces, and Arthrobacter. The most potent degrading NAHL strains were found among Bacillus spp. with about 60% isolation frequency, while four other genera were isolated at much lower frequencies (Arthrobacter 20%; Streptomyces 7%; Mesorhizobium 7% and Pseudomonas 6%).

Previews studies by Dong et al. (2000), Molina et al. (2003) and Park et al. (2003, 2005) revealed that Bacillus, Pseudomonas, Arthrobacter and Streptomyces isolates possess NAHL degrading properties. However, just one report was found to explain this trait in Mesorhizobium (Funami et al., 2005). All NAHL degrading isolates completely degrade NAHL in crude culture extract of EMPCC. 3-Oxo-C6HSL is signal molecule produced by P. carotovorum, which regulates production of virulence determinants such as extracellular enzymes was interrupted in standard quorum quenching experiments.

To set up quenching experiments we used Bacillus sp. strain DMS133, Arthrobacter sp. strain AM51 and Mesorhizobium sp. S5 as interfering agents, since during degradation assays these were demonstrated to be the most efficient NAHL-degrading activities. In quenching experiments, all tested isolates especially Bacillus sp. DMS133 inhibited pathogen growth rates and effectively reduced plant tissue maceration (Fig. 3). A possible approach in developing modern disease management is to target the QS genes of all putative bacterial NAHLs degraders were amplified by PCR using pA and pH primers as described above. DNA sequences were compared to those found in the data banks using the online FASTA search engine (http://www.ncbi.nlm.nih.gov). All the 16S rDNA gene sequences exhibited similarity between 85-100% with described genera. This is supported by preliminary identification of the isolates. Blast analysis indicated that nine of the isolates i.e. S6, S22, S27, AM35, AM38, EA60, EA73, EA85 and DMS133 belonged to Bacillus species. Within B. cereus group, the B. anthracis, B. cereus and B. thuringiensis were shown NAHL degrading activity with possible presence of NAHL-degrading aiiA gene (Dong et al., 2000). In this study, we isolated two strains of B. subtilis as NAHL degrading Bacillus species. B. subtilis have not been reported as an NAHL degrading species and does not belong to the B. cereus group. Isolate S5 showed 90% similarity to Mesorhizobium species, has been confirmed to degrade various NAHLs. The latter was already known to be able to degrade NAHLs. The two other identified genera with NAHL degrading activities were Arthrobacter (strains AM43, AM51 and S15) and one Streptomyces strain encoded as EA113 which already described as capable of NAHL degradation with long chain degradation preference (Park et al., 2005).
regulation system which controls the expression of major pathogenicity determinants (Dong et al., 2007; Sperandio, 2007). NAHL degrading rhizobacteria in this respect could potentially be then used for biological control NAHL producing plant pathogenic bacteria. Any success in their application will depend on population densities of the antagonists and the efficiency with which the NAHLS are inactivated (Jafra et al., 2006). Characterization of NAHL degrading bacteria from different sources, particularly from rhizosphere, will be helpful in identifying potential bacteria to be used as biocontrol agents. Dong et al. (2002), Molina et al. (2003), Jafra et al. (2006) and Morohoshi et al. (2009) reported a possibility for attenuating the virulence of *P. carotovorum* on potato introducing of quorum sensing interfering bacteria. These studies, as well as the data presented in this report, further support the promising strategy for the biocontrol and prevention of *P. carotovorum* infections through NAHL signal degradation.

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