Biological Control of Grapevine Crown Gall Caused by *Allorhizobiumvitis* using Bacterial Antagonists

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Abstract: The potential use of bacterial antagonists isolated from different area in Morocco for the biological control of grapevine crown gall caused by Allorhizobiumvitis was investigated. In vitro analyzing the activity of 90 bacterial isolates towards Allorhizobiumvitis strain S4 resulted in a selection of 26 biocontrol agents. The isolated were tested for their ability in vitro to inhibit the growth of the pathogen. Among these isolates 12 antagonists are efficient. Molecular identification of selected isolates, using rDNA 16S sequencing, show that the antagonists belong to different genera as Bacillus spp., Pantoea sp., Rahnella sp., Acinetobacter spp. and Enterobacter sp. Four antagonists were tested for their antagonistic effect in planta; they exhibited considerable inhibitory activity to reduce the incidence of galls in tomato and squash fruits. Rahnellaaquatilis and Pantoeaagglomerrans reduced the incidence of crown gall up to 100% both in tomato and squash fruits. Bacillus subtilis reduced the incidence of gall development to 75% in squash fruits and 60% in tomato. Whereas, Acinetobacter calcoaceticus reduced incidence of gall formation to 65% in squash fruits and 47% in tomato. Consequently, the treatment with bacterial antagonists may be used as an effectiveness alternative to control crown gall disease.

Keywords: Crown gall, grapevine, Allorhizobiumvitis, biological control, bacterial antagonists, Morocco

1. Introduction

Crown gall of grapevine is a bacterial disease caused by [39], Agrobacteriumvitis recently renamed Allorhizobiumvitis [37]- [38]. This disease presents a serious problem to grapevine production in several regions of the word [8]. A. vitis can survives systemically in the plant tissues until conditions become suitable for gall development [6] and, therefore, is disseminated in propagating plant material [30]- [50]. The tumorigenicity of A. vitisis determined by the presence of a set of oncogenes called the T-DNA that are carried on a large tumor-inducing (Ti) plasmid [51]. The T-DNA is transferred and integrated into the plant cell genome; the genes encode enzymes that stimulate production of plant hormones (auxin and cytokinin) that cause the proliferation of plant cells and development of tumors. The T-DNA also contains genes responsible for the biosynthesis of opines, specific carbon and nitrogen sources for A. vitis development [1]- [31].

The systemic survival of *A. vitis* in grapevine makes it difficult to controlthe disease. The older vines usually survive the infection, although they show stress symptoms, young vines that develop a tumor at their graft union often die [48]. The damage could threaten the harvest and grape quality in the absence of appropriate crop management [11]. Until now, the control of crown gall of grapevine is based on viticultural criteria as well as on the indexing and certification of propagation material [3]- [49]. Currently, there are no real effective chemical treatments to control the crown gall of grapevine. Generally, chemical options of this disease are limited to the use of disinfectants (sodium hypochlorite) copper or antibiotics [3]- [24]- [35]. However, these treatments can kill the bacteria on gall

surfaces, and fail to control the systemic survivalof *A. vitis* in the vascular tissue of the grapevine [7]- [10].

Recently, the biological control of grapevine crown gall has been the subject of several studies due to the absence of other effective means of control. The use of microorganisms to prevent the disease, offers an attractive alternative for the management of crown gall disease. Numerous bacterial antagonists with biological control activity have been evaluated against A. vitis [12]- [13]-[15]- [22]- [28]- [43], which can have direct (antibiosis, competition for target sites or nutrients) or indirect action (induced resistance in the host). Some of these antagonists are epiphytic; they colonize the rhizosphere and the root of host, while others cans also survive systematically in the hot and becomes endophytes [14]- [34]- [46]. The use of Agrobacterium radiobacter strain K-84 as a biological treatment was the first model used successfully against crown gall on several plant species caused by Agrobacterium tumefaciens [45]; however, this antagonist is not effective for preventing infections on grape caused by A. vitis [6]- [25]. Recent studies have demonstrated that the use of nonpathogenic strains of A. vitis inhibit growth of most tumorigenic strains of A. vitisin vitro and can also inhibit crown gall on grapevine in the greenhouse [21]-[32]- [47]- [53]. The main purpose of this study was to evaluate the practical potential of biological control of some bacterial antagonists isolated from Morocco against A. vitis strain S4 using in vitro and in planta.

2. Material and Methods

Bacterial strain and culture conditions

The bacterial strains used in this study are listed in Table 1. *A. vitis* strain S4 used in this study was cultivated on

MG medium [36](D-mannitol, 5g/L; L-glutamic acid, 2g/L; KH₂PO₄, 0.5g/L; NaCl, 0.2g/L; MgSO₄×7H₂O, 0.2g/L; Yeast extract, 0.5g/L; Agar, 15g/L; pH=7) and incubated, for 24 hours, at 28° C.

Antagonistic bacteria were selected among a collection of 90 isolates, which were isolated from different plants in different region in Morocco. They belong to the collection of laboratory of Phytobacteriology and Biological Control of the National Institute of Agronomic Research of Meknes. The selection of bacterial antagonist was based on the ability to inhibit growth of *A. vitis* in YPGA medium(yeast extract, 5g/L; peptone, 5g/L; glucose, 10g/L; agar, 15g/L). The strains were cultured at 28°C on YPGA and incubated, for 24 hours, at 28°C.

Antagonistic activity in vitro

A 100 μ l sample of the *A. vitis* strain S4 suspension (10⁷ CFU/mL) was inoculated, using the flooding method, on YPGA medium. Sterile filter paper discs (5mm diameter) was impregnated by bacterial cream of antagonist and placed in Petri dishes, either directly onto the center of the culture medium. The plates were incubated in the upright position at 28°C for 24 hours. The filter soaked with 2 μ L of sterile distilled water was served as negative control and filter impregnate with streptomycin (32mg/L) was served as positive control. After incubation, the inhibition zone around each disc was measured and the percentage inhibition was calculated using the following formula [41]:

 $Percent inhibition (\%) = \left[\frac{(Rate without inhibitor - Race with inhibitor)}{Rate whitout inhibitor}\right] \times 100$

Molecular identification of antagonistic bacteria

The identification of bacterial antagonists was made for 12 antagonistic bacteria, presenting a height antibacterial activity against *A. vitis* strain S4, using conserved 16S rRNA gene for the detection and identification of bacteria. The DNA extraction was made using alkaline method [42]. Pure culture genomic DNAs were extracted from bacteria grown overnight at 28°C in YPGA. One colony of each bacteria isolated was mixed with 10µl of (20 mMNaOH) and incubated at 37°C for 5 minutes. The bacterialyses cells were stored at 4°C until they use.

Amplification was carried with F809pA primers (AGAGTTTGATCCTGGCTCAG) F810pH and (AAGGAGGTGATCCAGCCGCA). Standard PCR was carried out in a 60µl reaction volume containing 38.6µL H₂O, 6µl (2mM) DNTPs, 1.2µl (2mM) MgCl₂, 3µl DMSO (Dimethyl sulfoxide), 1µl (10 µM) of each primer, 0.2µl Taq DNA polymerase (Invitrogen, France) and 3µl of lyses cells. The PCR was performed using the following program: initial denaturation at 94°C for 5min, followed by 35 cycles of denaturation at 94°C for 1min, annealing at 55°C for 1min and extension at 72°C for 1min, followed by an additional extension at 72°C for 5min.

Electrophoresis was performed in 1% agarose gel. The gel was stained with ethidium bromide. Fragments were

visualized with an ultraviolet (UV) transilluminator, and the gel was photographed. The 16S gene of each isolates was sequenced (GenoScreen Lille-France) and analyzed using NCBI-BLAST software [2].

Biocontrol activity in planta

Four isolates that yielded the greatest percentage of inhibition for growth of *A. vitis* stain S4 were selected to demonstrate its biocontrol activity *in planta*as a preventive treatment against tumor development. These isolates were examined for their ability to suppress gall formation by *A. vitis* in tomato (*Solanum lycopersicum* L.) and summer squash fruits (Cucurbita pepo cv. Eskandarany). Each test plant was inoculated with one of the biocontrol agents.

In the case of tomato, the inoculation was made by 10 μ l of specific antagonistic suspension (10⁷CFU/mL) of 24 hours bacterial culture in stem internodes of tomato 2-3 weeks after transplanting. After the liquid was absorbed by the plant tissue, the wounded sites were wrapped in Parafilms. After 24 hours, each site of inoculation was rewounded and 10 μ l of *A. vitis* strain S4 suspension was introduced in each site. Non-treated plants inoculated with *A. vitis* S4 or with sterile distilled water were used as a positive control and negative control, respectively. After the suspension was absorbed into the wound, the stem was again wrapped in Parafilms. The inoculated plants were maintained in greenhouse at 27°C during 3-4 weeks. Number and size of formed galls were recorded.

In the case of squash fruits, uniform fruits (10 to 15 cm of length and 3 to 5 cm of width) were stabbed by toothpick to make holes at five sites distributed over two rows (5 sites/row) per fruit. The inoculation was performed in the same manner as in tomato leaves. The treated fruits were maintained in plastic containers with transparent plastic covers and kept at 28°C. The presence or absence of tumors was visible 7 weeks or 10 days after inoculation and the number and size of formed galls were recorded.

Statistical Analysis

The significant effect of bacterial antagonists on growth inhibition of *A. vitis* was evaluated by Analysis of variance (ANOVA1) (factor: treatment), performed with the SPSS 20 statistical software ((IBM Corporation, Somers, NY, USA). The arcsin of the inhibition percentage was used for statistical analysis and was calculated using the formula $\text{Arcsin}=\sqrt{(\%I/100)}$, where %I is the rate of bacterial growth inhibition.

3. Results

Antagonistic activity in vitro

Among 90 isolates, many isolates showed antagonistic activity toward *A. vitis* strain S4 in variable degree. Among these isolates, only 26 isolates exhibited considerable inhibitory activity (Figure 1). The mean values of percent of inhibition resulted from these isolated fluctuated between 13.3 and 39.8%. The greatest inhibition resulted from isolate 2515-3 (39.8%) flowed by isolate

2332-A1 (32.2%) (Figure 2). The isolates 2626-5, 2510-8, 2627-1, 2510-9, 2027-1, 2546-4, 2066-7, 2328-B5, 2021-12 and 2021-2 showed moderately antagonistic reaction indicated by the percent of inhibition values (25.5, 16.6, 27.7, 24.6, 27.7, 27.7, 27.7, 27.9, 22.22, 20% respectively).

Molecular identification of antagonist

Further molecular analysis was carried out using universal primers F809PA and F810PH targeted gene 16SrDNA of 1477 bp (Figure 3). Analysis of the 16SrDNA sequence, by BLAST-NCBI, of 12 bacterial antagonist perentinga high efficacy against *A. vitis* S4 and originating from different samples and locations in Moroccoare shown in Table 2. According to the sequencing results of 16SrDNA, the bacterial antagonists belongs to different species from different genus: *Bacillus*spp., *Pantoeas*pp., *Rahnella*spp., *Acinetobacters*pp. and *Enterobacters*pp. The antagonists strain belong to *Pantoea* and *Rahnella* genus possessed a 16S rDNA sequence with \geq 99% similarity to that of genus members. The *Enterobacter* and *Acinetobacter* antagonists exhibited \geq 97% 16S rDNA similarity whit this genus.

Biocontrol activity in planta

Inoculation of wounded sites on tomato and squash fruits with antagonistic bacterial isolates, using as a prevent treatment, provide significant reduction in incidence and size of galls formed in response to subsequent inoculation with A. vitis S4 comparatively with 100% gall incidence resulted with positive control (inoculation with A. vitis S4 alone). There was significant difference between the tested antagonists in the prevention of galls formation on the different tested plants (Figure 4, Figure 5). The antagonists (Rahnellaaquatilis) 2332-A1 2066-7 and (Pantoeaagglomerans) reduced the incidence of crown gall up to 100% in tomato and squash fruits. The isolate 2515-3 (Bacillus subtilis) reduced incidence of crown gall to 75% in squash fruits and 60% in tomato. Isolate 2328-B5 (Acinetobacter calcoaceticus) reduced incidence of gall formation to 65% in squash fruits and 47% in tomato.

4. Discussion

In this study, the efficacy of treatment with bacterial antagonists was demonstrated by the reduction in vitro as well as in planta. From 90 tested microbial antagonists, only 26 isolates exhibited an antibacterial activity in vitro against A. vitis S4. The idea provided to select effective biocontrol strain, among the collection, capable to control A. vitisin vitro and in planta. Numerous studies were conducted to evaluate the antibacterial activity of some microbial antagonists and their potential for use in biocontrol programs for the management of grapevine crown gall [52]. The nonpathogenic strains of A. vitis were the first model has been studied as a biological control agent of grapevine crown gall. Numerous strains of A. vitis were effective in vitro and in vivo against tumorigenic strains. A. vitis strain F2/5 was the most effective strain capable to inhibit the growth of pathogenic strains in vitro and galls development in grapevine [4]- [9]- [10]- [26]. The mechanism of action of A. vitis strain F2/5 is the

competition for attachment sites on grape and antibiosis mechanisms by producingantibiotic (agrocin) [9].

Among the screened isolates in this study, 12 antagonistic bacteria, with strong antibacterial activity in vitro against A. vitis S4, were selected and identified in the genera Bacillus (B. subtilis and B. cereus), Pantoea (P. agglomerans), Rahnella (R. aquatilis), Acinetobacter (A. calcoaceticusand A. venetianus) and Enterobacter (E.ludwigii). Moreover, the result of the present study demonstrates that the bacterial antagonists tested can inhibit the growth of A. vitisin vitro. However, B. subtilis (2515-3) is the most effective biocontrol agent in vitrofollowed by R. aquatilis (2332-A1) and P. agglomerans (2066-7). The efficacy of these species has been documented for antibacterial activities in vitro against many pathogens include A. vitis. In the study work of Sholberget al. [43], they demonstrate that two species of Bacillus spp. (EN63-1 and E71-1) can inhibit bacterial growth of A. vitis in vitro. The efficacy of R. aquatilis was also evaluated against A. vitis by Bell et al. [5]; Chen et al. [12]- [13]; when they showed that this antagonist can exhibit a high antibacterial activity against A. vitis strains in vitro and in planta. In the research work of Kenneth et al. [28], they show that the P. agglomerans inhibit the growth of A. vitisin vitro and can inhibit galls development in planta.

Some bacterial antagonists are known to have antibacterial activities in vitro but the biocontrol effectiveness may not be expressed under in planta conditions [23]. In the present research, the selected antagonistic isolates proved to be efficient in vitro and under in planta conditions in variable degree. R. aquatilis (2332-A1) and P. agglomerans (2066-7) are the most effective antagonists capable to exhibit considerable inhibitory activity toward gall formation both in tomato and squash fruits (100% of reduction). This correlation between in vitro and in planta results have been documented in the study of Bell et al. [5], Chen et al. [12]- [13], Tolba and Solimane [52] and Gupta et al. [20]. Interestingly, the antagonistic isolate B. subtilis (2515-3), which present the most effective antagonist in vitro, reduce the galls development only to 60% in tomato and 75% in squash fruits. This lack of correlation between in vitro and in planta tests, in the control of A. vitis, has been documented in other studies with other antagonists [4]- [29]- [52]. For the isolate A.calcoaceticus (2328-B5) reduced the incidence of crown gall to 65% in squash fruits and 47% in tomato. This experiment indicated that the different in the performance of antagonistic bacteria has been attributed to variability in the physical and chemical properties within the niches occupied by biocontrol agents and by the host, which affect the colonization and efficacy of biocontrol agent [40].

Biological control using antagonistic bacteria result from many different types of interactions between organisms. The information about the mechanism of action for most of the antagonists is still incomplete due to difficulties in analyzing the complex interactions between host, pathogen, antagonist and other microorganisms present [46]. Several biocontrol mechanisms have been described including antibiosis, competition for nutrients and space, induction of mechanisms of resistance in the host plant, quorum quenching and direct interaction between the antagonist and the pathogen including parasitism. Actually, most biocontrol agents not only use one mechanism of biocontrol, but disease control is the result of a combination of several mechanisms [18]- [44]. Bacterial antagonists members of the genus Bacillus were reported, in many studies, to exhibit an antagonistic activity by producing a wide range of secondary metabolites such as antibiotics (iturin and gramicidin) [16]- [19], non-volatile and volatile compounds and lytic enzymes [17]- [43]. R. aquatilis was reported to synthesis a secondary compounds (antibiotic) with wide range of action against bacteria and fungus [33]. P. agglomerans have a preventive effect by competition mechanism to acquire nutrients from the environment than the pathogen; is the most antagonist presenting a higher competitive ability used against many pathogenic bacteria and fungus [27].

5. Conclusion

In conclusion, results obtained in the present work show that the bacterial isolates identified may be considered as potential sources of bioactive metabolites and an important alternative to control grapevine crown gall disease. They provide a crop protection with a low environment risk associated. Future studies are recommended to develop a mass protection method of the bacterial antagonists, to find the appropriate formulation that allow to increase biocontrol activity and ensure its stability and to develop a bacterial pesticides used in the control of crown gall of grapevine.

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References

- Akiyoshi, D.E., Klee, H., Amasino, R.M., Nester, E.W., Gordon, M.P. (1984). T-DNA of *Agrobacterium tumefaciens* encodes an enzyme of cytokinin biosynthesis. *ProcNatlAcadSci USA*, 81:5994-5998.
- [2] Altschul, S.F., Madden, T.L., Schaffer, A.A., Zhang, J., Zhang, Z., Miller, W., Lipman, D.J. (1997). Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids.

- [3] Armijo, G., Schlechter, R., Agurto, M., Munoz, D., Nunez, C., Arce-Johnson, P. (2016). Grapevine pathogenic microorganisms: understanding infection strategies and host response scenarios. *Front Plant Sci*, doi: 10.3389/fpls.2016.00382.
- [4] Bazzi, C., Alexandrova, M., Stefani, E., Anaclerio., Burr, T.J. (1999). Biological control of *Agrobacterium vitis*using non-tumorigenic agrobacteria. *Vitis*, 38 (1):31-35.
- [5] Bell, C.R., Dickie, G.A., Chan, J.W.Y.F. (1995). Variable response of bacteria isolated from grapevine xylem to control grape crown gall disease *in planta*. *Am J EnolVitic*, 46: 499–508.
- [6] Burr, T.J., Bazzi, C., Sul, S., Otten, L. (1998). Crown gall of grape: biology of *Agrobacterium vitis* and the development of disease control strategies. *Plant Dis*, 82 (12): 1288-1297.
- [7] Burr, T.J., Reid, C.L., Splittstoesser, D.F., Yoshimura, M. (1996). Effect of heat treatment on grape bud mortality and survival of *Agrobacterium vitis in vitro* and in dormant grape cuttings. *J Enol Vitic*, 47(2): 119-123.
- [8] Burr, T.J., Reid, C.L. (1993). Biological control of grape crown gall with nontumorigenic Agrobacterium vitis F2/5. Am J EnolViticult, 45:213-219
- [9] Burr, T.J., Reid, C.L., Taglicti, E., Bazzi, C., Süle, S. (1997). Biological control of grape crown gall by strain F2/5 is not associated with agrocin production or competition for attachment site on grape cells. *Phytopathology*, 87:706-711.
- [10] Burr, T.J., Otten, L. (1999). Crown gall of grape: biology and disease management. AnnuRev Phytopathol, 37:53–80.
- [11]Burr, T.J. (2004). Grape crown gall biology and strategies for control. *Foundation Plant Services*, *Grape Program Newsletter*, *University of California-Davis*, pp16-18.
- [12] Chen, F., Guo, Y.B., Wang, J.H., Li, J.Y., Wang, H.M. (2007). Biological control of grape gall by *Rahnellaaquatilis* HX2. *Plant Dis*, 91:957-963.
- [13] Chen, X.H., Scholz, R., Borriss, M., Junge, H., Mogel, G., Kunz, S., Borriss, R. (2009). Difficidin and bacilysin produced by plant-associated *Bacillus amyloliquefaciens* are efficient in controlling fire blight disease. *J Biotechnol*, 140:38-44.
- [14] Dunne, C., Cronin, D., MoenneLoccoz, Y., O'Gara, F., Duffy, B., Rosenberger U., Defago, G. (1998).
 Biological control of phytopathogens by phloroglucinol and hydrolytic enzyme producing bacterial inoculants. *Molecular approaches in biological control. Delemont, Switzerland, 15–18* September 1997. Bulletin OILB SROP, 21: 19–25.
- [15] Eastwell, K., Sholberg, P., Sayler, R.(2006). Characterizing potential bacterial biocontrol agents for suppression of *Rhizobium vitis*, causal agent of crown gall disease in grapevines. *Crop Protect*, 25:1191– 1200.
- [16] Edwards, S.G., Seddon, B. (2001). Mode of antagonism of *Brevibacillus brevis* against *Botrytis cinereain vitro*. *JApplMicrobiol*, 91:652-659.
- [17] Frandberg, E., Schnurer, J. (1998). Antifungal activity of chitinolytic bacteria isolated from airtight stored cereal grain. *CanJMicrobiol*, 44:121-127.

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- [18] Gonzalez, MC.M. (2007). Characterization and mechanism of action of the biological control *Pantoeaagglomerans* EPSI25. *Doctoral thesis*.
- [19] Gueldner, R.C., Reilly, C.C., Pusey, P.L., Costello, C.E., Arrendale, R.F., Cox, R.H., Himmelsbach, D.S., Grumley, F.G., Cutler, H.G. (1988). Isolation and identification of iturins as antifungal peptides in biological control of peach brown rot with *Bacillus subtilis*. *JAgriFoodChem*, 36:366-370.
- [20] Gupta, A.K., Kishore, K., Bhardwaj, S.S., Aman, T., Sapna, D., Jarial, R.S., Chhaya, S., Singh, K.P., Srivastava, D.K., Rup, L.(2010). Biological control of crown gall on peach and cherry rootstock colt by native Agrobacterium radiobacter isolates. Open Horticult J, 3:1-10.
- [21] Hao, G., Zhang, H., Zheng, D., Burr, T.J. (2005). LuxR homolog avhR in Agrobacterium vitis affects the development of a grape-specific necrosis and a tobacco hypersensitive response. J Bacterio, 187:185-192.
- [22] Herlache, T.C., Triplett, E.W. (2002). Expression of a crown gall biological control phenotype in an avirulent strain of *Agrobacterium vitis* by addition of the trifolitoxin production and resistance genes. *BMC Biotechnology*, doi: 10.1186/1472-6750-2-2
- [23] Inam-ul-Haq, M., Javed, M., Ahmad, R., Rehman, A.(2003). Evaluation of different strains of *Pseudomonas fluorescens* for the biocontrol of fusarium wilt of chick pea. *Pakistan J Plant Pathol*, 2:65-74.
- [24] Kawaguchi, A. (2009). Studies on the diagnosis and biological control of grapevine crown gall and phylogenetic analysis of tumorigenic *Rhizobium vitis*. *J Gen Plant Pathol*, 75:462.
- [25] Kawaguchi, A., Inoue, K., Nasu, H. (2005). Inhibition of crown gall formation by *Agrobacterium radiobacter*biovar 3 strains isolated from grapevine. J Gen Plant Pathol, 71:422-430.
- [26] Kawaguchi, A., Sawada, H., Ichinose, Y. (2008). Phylogenetic and serological analyses reveal genetic diversity of *Agrobacterium vitis* strains in Japan. *Plant Pathol*, 57:747-753.
- [27] Kempf, H.J., Wolf, G. (1989). Erwiniaherbicola as a biocontrol agent of Fusarium culmorum and Puccinia recondite F SpTritici on wheat. Phytopathology, 79:990-994.
- [28] Kenneth, C.E., Peter, L.S., Ronald, J.S. (2006). Characterizing potential bacterial biocontrol agents for suppression of *Rhizobium vitis*, causal agent of crown gall disease in grapevines. *Crop Protection*, doi: 10.1016/j.cropro.2006.03.004
- [29] Kerr, A., Htay, K. (1974). Biological control of crown gall through bacteriocin production. *Physiol Plant Pathol*, 4:37-44.
- [30] Kuzmanovic, N., Gasic, K., Ivanovic, M., Prokic, A., Obradovic, A. (2012). Identification of *Agrobacterium vitis* as a causal agent of grapevine crown gall in Serbia. *Arch Biol Sci*, 64(4): 1487-1494.
- [31] Lacroix, B., Citovsky, V. (2013). Crown gall tumors. Brenner's Encyclopedia of Genetics, 2(2): 236-239.
- [32] Li, J.Y., Wang, J.H., Wang, H.M.(2009). Modeofaction of the antibacterial compound Ar26 produced by Agrobacteriumvitis strain E26 with

activity against *A.tumefaciens*, *A.rhizogenes* and *A.vitis*. JPhytopathol, 157:159-165.

- [33] Li, L., Ziwei, J., Lauren, H., Wu, W., Guo, Y. (2014). Distruption of gene *pqqA* or *pqqB* reduces plant growth promotion activity and biocontrol of crown gall disease by *Rahnellaaquatilis* HX2. *Plos one*, 9(12):e115010, doi:10.1371/journal.pone.0115010
- [34] Lugtenberg, B., Chin-A-Woeng, T., Bloemberg, G. (2002). Microbe-plant interactions: principles and mechanisms. *Antonie van Leeuwenhoek*, 81:373-383.
- [35] McManus, P.S., Stockwell, V.O., Sundin, G.W., Jones, A.L. (2002). Antibiotic use in plant agriculture. *Annu Rev Phytopathol*, 40: 443-465.
- [36] Moore, L.W., Kado, C.I. Bouzar, H. (1988). Agrobacterium. In laboratory guide for identification of plant pathogenic bacteria: 16-36. Nws.c.h.a.a.d., aps press, saint paul, minnesota.
- [37] Mousavi, S.A., Osterman, J., Wahlberg, N., Nesme, X., Lavire, C., Vial, L., Paulin, L., de Lajudie, P. et al. (2014). Phylogeny of the *Rhizobium-Allorhizobium-Agrobacterium* clade supports the delineation of *Neorhizobium* gen. nov. *SystApplMicrobiol*, 37:208-215.
- [38] Mousavi, S.A., Willems, A., Nesme, X., de Lajudie, P., Lindstrom, K. (2015). Revised phylogeny of Rhizobiaceae: proposal of the delineation of *Pararhizobium* gen. nov., and 13 new species combinations. *SystApplMicrobiol*, 38:84-90.
- [39] Ophel, K., Kerr, A. (1990). Agrobacterium vitis sp. nov.for strains of Agrobacterium biovar 3 from grapevines. Int J Systbacterial, 40: 236-241.
- [40] Ryan, A.D., Kinkel, L.L., Schottel, J.L.(2004). Effect of pathogen isolate potato cultivar and antagonist strain on potato scab severity and biological control. *Biocontrol SciTechnol*, 14:301-311.
- [41] Schultz, D.L. (2006). Biology 155 General Biology I Laboratory Supplement, pp78.
- [42] Shams, M., Campillo, T., Lavire, C., Muller, D., Nesme, X., Vial, L. (2012). Rapid and efficient methods to isolate, type strains and determine species of *Agrobacterium* spp. in pure culture and complex environments. *In* Dr. Jose C. Jimenez-Lopez (ed.)-Biochemical Testing. *In Tech*, doi: 10.5772/2250.
- [43] Sholberg, P.L., Marchi, A., Bechard, J.(1995). Biocontrol of postharvest diseases of apple using *Bacillus* spp. isolated from stored apples. *Can J Microbiol*, 41:247–252.
- [44] Siddiqui, Z.A., Mahmood, I.(1999). Role of bacteria in the management of plant parasitic nematodes. *Rev BioresourTechnol.* 69 :167–179.
- [45] Smith, E.F., Townsend, C.O. (1907). A plant-tumor of bacterial origin. *Science* (newyork), 25: 671-673.
- [46] Spadaro, D., Gullino, M.L. (2004). State of the art and future prospects of the biological control of postharvest fruit disease. *Int J Food Microbiol*, 91:185-194.
- [47] Staphorst, J.L., VanZyl, F.G.H., Strjidom, B.W., Üroenewold, Z.E.(1985).Agrocin-producing pathogenic and nonpathogenic biotype-3 strains of *Agrobacterium tumefaciens* active against biotype-3 pathogens. *CurrMicrobiol*, 12: 45-52.
- [48] Sul, S., Burr, T.J. (1998). The influence of rootstock resistance to crown gall (*Agrobacterium spp.*) on the

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susceptibility of scions in grape vine cultivars. *Plant Pathol*, 47: 84-88.

- [49] Szegedi, E., Nemeth, 1. (1996).Investigation of grape shoots for Agrobacterium vitis. Növenyvede1em, 32: 605-609.
- [50] Tarbah, F.A., Goodman, R.N. (1986). Rapid detection of *Agrobacterium tumefaciens* in grapevine propagating material and the basis for an efficient indexing system. *Plant Dis*, 70:566-568.
- [51] Thomashow, L.S., Reeves, S., Thomashow, M.F. (1984). Crown gall oncogenesis: evidence that a T-

DNA gene from the *Agrobacterium* Ti plasmid pTiA6 encodes an enzyme that catalyzes synthesis of indoleacetic acid. *Proc Natl AcadSci*, 81: 5071-5075.

- [52] Tolba, I.H., Soliman, M.A.(2013). Efficacy of native antagonistic bacterial isolates in biological control of crown gall disease in Egypt. *Ann AgriScience*, 58(1): 43-49.
- [53] Wang, H.M., Sun, Y.L., Wang, J.H. (2002). Preliminary study of biological control agent *Agrobacterium vitis* strain E26 on some ecological factors. *SciAgricul Sin*, 35:38-41

Strains	Relevant characteristics	Sampling location
Allorhizobium vitis S4	Isolated from black raspberry	Hungary
2066-7	Isolated from Olivier (Pichouline) olive knot	Taounat
2072-2	Isolated from onion	Ifrane
2021-12	Isolated from compost	Agadir
2022-18	Isolated from Tomato	Casablanca
2546-3	Isolated from strawberry (verticillium)	Laïche
2544-3	Isolated from strawberry (verticillium)	Laïche
2332-A	Isolated from apple (crown)	El Hajeb
2627-1	Isolated from apple (crown)	El Hajeb
2026-2	Germ	INRA Meknes
2015-3	Isolated from compost	Meknes
2510-8	Isolated from Olivier (Pichouline) olive knot	Meknes
2546-4	Isolated from strawberry (verticillium)	Laïche
2021-9	Isolated from compost	Agadir
2015-8	Isolated from compost	Meknes
2328-B5	Isolated from apple (crown)	Fez
2261-2	Isolated from gravine (Muscat)	El Hajeb
2072-22	Isolated from compost	INRA Meknes
2515-3	Isolated from apple	Imouzzar Kandar
2510-9	Isolated from Olivier (Pichouline) olive knot	Meknes
2026-4	Isolated from compost	Meknes
2021-20	Isolated from compost	Agadir
2027-1	Isolated from compost	Meknes
2626-5	Isolated from apple	El Hajeb
2021-2	Isolated from compost	Agadir
2025-6	Isolated from compost	Meknes
2015-7	Isolated from compost	Meknes

Table 1: strains used in this study

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Table 2: 16 rDNA results of 12 bacterial antagonists tested in this study			
Code	Species	GenBank accession No	
2515-3	Bacillus subtilis	KJ592619.2	
2626-5	Pantoea agglomerans	KJ781904.1	
2510-8	Pantoea sp	HQ396801.1	
2332-A1	Rahnella aquatilis	KM241863.1	
2627-1	Acinetobacter calcoaceticus	KP170504.1	
2510-9	Pantoea ananatis	KM977993.1	
2027-1	Bacillus cereus	KR493006.1	
2546-4	Enterobacter ludwiqii	LC015543.1	
2066-7	Pantoea agglomerans	KJ781904.1	
2328-B5	Acinetobacter calcoaceticus	KP170504.1	
2021-12	Acinetobacter venetianus	KP009554.1	
2021-2	Bacillus cereus	KR493006.1	

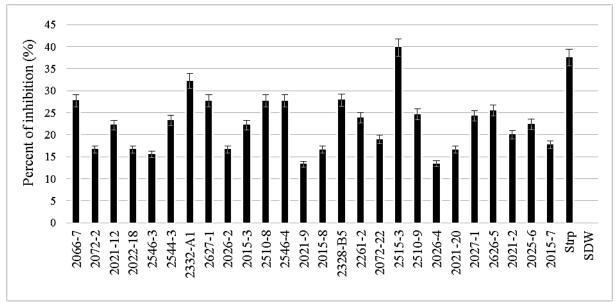


Figure 1: In vitro percent of inhibition caused by bacterial antagonists against Allorhizobiumvitis (strain S4). Strep: streptomycin antibiotic, SDW: sterile distilled water

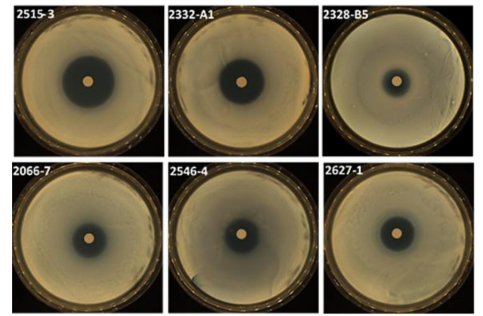


Figure 2: Inhibition zones resulted from challenge of bacterial isolates toward Allorhizobiumvitis (strain S4)

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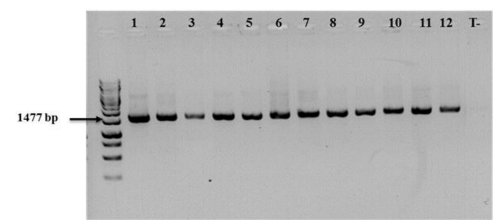


Figure 3: Electrophoritic profile of bacterial antagonits: (1) 2515-3. (2) 2626-5. (3) 2510-8. (4) 2332-A1. (5) 2627-1. (6) 2510-9. (7) 2027-1. (8) 2546-4. (9) 2066-7. (10) 2328-B5. (11) 2021-12. (12) 2021-2. (T-) negative control

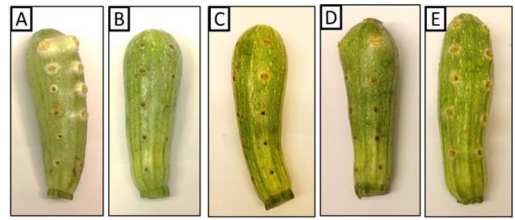


Figure 4: Effect of four antagonistic bacterial isolates on incidence and size of galls induced by *Allorhizobiumvitiss*train S4 in squash fruits. (A) positive control (inoculation with *A. vitis* S4), (B to E) preventive treatment with bacterial antagonists; (B) 2332-A1 (*R. aquatilis*), (C) 2066-7 (*P. agglomerans*), (D) 2515-3 (*B. subtilis*) and (E) 23228-B5 (*A. calcoaceticus*)



Figure 5: Effect of four antagonistic bacterial isolates on incidence and size of galls induced by *Allorhizobiumvitis*strain S4 in tomato plant. (A) positive control (inoculation with *A. vitis* S4), (B to E) preventive treatment with bacterial antagonists; (B) 2332-A1 (*R. aquatilis*), (C) 2066-7 (*P. agglomerans*), (D) 2515-3 (*B. subtilis*) and (E) 23228-B5 (*A. calcoaceticus*)

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