# **Short Paper**

# First detection of *Pantoea ananatis*, the causal agent of bacterial center rot of onion in Morocco

El Hassan Achbani<sup>1</sup>, Hamid Mazouz<sup>2</sup>, Abdellatif Benbouazza<sup>1</sup> and Soumia Sadik<sup>1, 2\*</sup>

1. Laboratory of Plant Protection, URPP- INRA-Meknes, Morocco.

2. Laboratory of Plant Biotechnology and Molecular Biology, Faculty of Sciences, Meknes, Morocco.

Abstract: *Pantoea ananatis* is an important pathogen causing bacterial center rot in onion. Isolated strains from onion soft rot bulbs based on morphological characteristics were tested for hypersensitivity reaction on tobacco. Pathogenic strains underwent some biochemical tests which detected the presence of *P. ananatis*, based on RNA16S sequencing, this result was confirmed. Symptoms were reproduced by inoculating this species to onion leaves and bulbs. This study reports for the first time the presence of *P. ananatis* bacterium in Morocco causing the bacterial center rot in onion.

Keywords: Allium cepa, bacterial center rot, Morocco, Onion, Pantoea ananatis

# Introduction

In Morocco, the onion crop (*Allium cepa* L.) has a high economical importance, representing 11% of nationally produced vegetable crops (Anonymous, 2011). Therefore, detection and control of diseases of this plant is important. Thus onion crop is exposed to many physiological and phytosanitary constraints (Conn *et al.*, 2012), among which is the bacterium center rot caused by *Pantoea ananatis*.

The enterobacterium *P. ananatis* is an ecologically versatile species. It has been found in the environment, as plant epiphyte and endophyte, emerging phytopathogen, and as a presumptive, opportunistic human pathogen (De Maayer *et al.*, 2012).

*Pantoea ananatis* causes disease symptoms in a wide range of economically important agricultural crops and forest trees all over the world. It has been reported to induce diseases in onion (Gitaitis and Gay, 1997; Schwartz and Otto, 2000), Eucalyptus

(Coutinho *et al.*, 2002), corn (Paccola- Meirelles *et al.*, 2001), melons (Bruton *et al.*, 1986; Wells *et al.*, 1987), Sudan grass (Azad *et al.*, 2000), rice (Azegami *et al.*, 1983), maize (Pérez-y-Terrón *et al.*, 2009) and pineapple (Serrano, 1928).

The onion seed lot associated with the first outbreak of center rot in Georgua, USA (Gitaitis and Gay, 1997), was reportedly produced in South Africa and Walcott *et al.* (2002)) suggested that the center rot pathogen was possibly introduced on infested seed lots. *P. ananatis* is both seed-borne and seed-transmitted in Sudan grass (Azad *et al.*, 2000), rice (Tabei *et al.*, 1988) and onions (Walcott *et al.*, 2002).

The disease affects the center leaves of onions, which become water-soaked, soft, and bleached white as the rot progresses. Advanced stages of the disease result in complete wilting and bleaching of all leaves (Mark *et al.*, 2002; Walcott *et al.*, 2002). Walcott *et al.*, 2002, reported natural infestation and transmission of *P. ananatis* in onion seed and implied that seed was the primary source of inoculum. The main objective of this work was to isolate and identify the causal agent of the bacterial soft rot on onion bulbs in Morocco using biochemical, molecular and pathogenicity tests.

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<sup>\*</sup> Corresponding author, e-mail: sadik.soumia@hotmail.com Received: 24 November 2014, Accepted: 03 November 2015 Published online: 24 November 2015

#### **Materials and Methods**

#### Isolation

Onion bulbs showing soft rot symptoms (Fig. 1) were sampled on 27 March 2013 from vegetables storage areas at Elhajjeb (N33°41.45, W5°22.00) (Meknes-Tafilalt, Morocco).

Onion bulbs showing center rot were used for isolation. These were washed with tap water and cut longitudinally. The diseased scale tissues were cut into 5 mm cubes using sterilized surgical blade. Three pieces of onion scale were ground in 1 ml of distilled water using a mortar and pestle. The suspension was streaked onto YPGA medium (yeast extract, 5 g; peptone, 5 g; glucose, 10 g; of agar, 18 g; and distilled water, 1000 ml). Six bacterial isolates were identified by biochemical and physiological tests including Gram stain; Levan production, Oxidase reaction, Arginine dihydrolase [ADH] and Tobacco hypersensitivity test, Catalase, Gram test (KOH), and API 20 gallery. All tests were repeated at least twice (Schaad et al., 2001).

# Pathogenicity test

Cultures of bacterial isolates obtained from onion bulbs, labeled 2276-1, 2276-2, 2276-3, 2276-4, 2276-5 and 2276-6 were used in a greenhouse pathogenicity experiment. In vitro, onion bulbs were inoculated by a suspension  $(10^8 \text{ CFU}/\text{ml})$  of 24-h bacterial culture; the vegetable material was incubated in moist chamber at a temperature of 30 °C, six days after incubation symptoms were observed. All tests were repeated at last twice.

# **PCR** amplification

To confirm the biochemical results, a molecular identification by sequencing was performed. The DNA extraction was made using GenElute Mammalian Genomic Kit (Anonymous, 2010). Quantification of DNA was performed using spectrophotometry. Amplification was carried with primers Fd1 (CAGAGTTTGATCCTGGCTCAG) and RP2 (AGAGTTTGATCCTGGCTCAG) (Klouche et al., 2007; Parola et al., 2002) at a PCR kit (Invitrogen). The PCR was accomplished in a total volume of 25 µl of the following reaction mixture: 2.5 µl 10X Buffer, 2 µl dNTP (10 mm),

 $0.125 \mu$ l of each primer (100  $\mu$ M), 0.75  $\mu$ l MgCl2 (50 mM), 0.2  $\mu$ l Taq (5 units /  $\mu$ l) and 5  $\mu$ l DNA. The PCR was performed using the following protocol: initial denaturation at 96 °C for 4 min, followed by 35 cycles of denaturation at 96 °C for 0.1 min, annealing at 52 °C for 0.4 min, and extension at 72 °C for 2 min, followed by an additional extension at 72 °C for 4 min.

Electrophoresis was performed in 1.5% agarose gel. Purification of PCR products was carried using the enzyme EXO-SAT according to the following schedule: 37 °C for 15 min followed by 80 °C for 15 min.



Figure 1 Onion bulbs showing soft rot.

# 16S DNA sequencing and sequence analysis

Selected PCR fragments, amplified from the isolates tested for pathogenicity, were sequenced in both strands, with the pA and pH' primers, using the BigDye terminator cycle sequencing ready reaction FS kit and an ABI PRISM 3700 DNA sequencer (both from Applied Biosystems Div., Foster City, Calif). So the sequencing of amplification product was carried out in a total volume of 10 µl of the following reaction mixture: 1 µl BigDye, 3 µl sequencing buffer x5, x µl primer (3.2-5 pmol), (0.75-1.5) µl DNA matrice, (2.5-3.25) µl H2O (MiliQ). The sequencing was performed using the following protocol: initial denaturation at 96 °C for 1 min, followed by 25 cycles of denaturation at 96 °C for 10 s, annealing at 50 °C for 5 s, and an extension at 60 °C for 4 min, followed by an additional extension at 72 °C for 4 min.

Reading of sequencing results was done using NCBI-BLAST software (Altschul et al., 1997).

# **Results and Discussion**

Bacterial colonies, which were consistently isolated from infected samples, were yellow and opaque on YPGA medium. All six representative strains isolated were Circular-shaped, Gramnegative, oxidase negative and catalase and arginine dihydrolase positive. The bacterial strains were able to induce hypersensitive reaction on tobacco leaves. All strains were capable of producing acid from sorbitol, sucrose, glucose, arabinose, melibiose but not able to utilize lysine (LDC), ornithine (ODC). thiosulfate (H<sub>2</sub>S), Urea (URE), sodium pyruvate (VP), inositol (INO), gelatin (GEL), amygdaline (AMY). However, the strains showed ability to utilized. indole ONPG (beta-(IND), galactosidase enzyme), CIT (citrate), TDA (tryptophan deaminase), MAN (mannitol) and RHA (rhamnose) (Table 1).

Table 1 Pantoea ananatis strains (2276-1, 2276-2, 2276-3, 2276-4, 2276-5 and 2276-6) behavior on gallery API20 and classic biochemical tests.

Tests	2276-1	2276-2	2276-3	2276-4	2276-5	2276-6	P. ananatis reference strain
IND	(+)	(+)	(+)	(+)	(+)	(+)	(+)
VP	(-)	(-)	(-)	(-)	(-)	(-)	(-)
GEL	(-)	(-)	(-)	(-)	(-)	(-)	(-)
Glucose	(+)	(+)	(+)	(+)	(+)	(+)	(+)
MAN	(+)	(+)	(+)	(+)	(+)	(+)	(+)
INO	(-)	(-)	(-)	(-)	(-)	(-)	(+)
Sorbitole	(+)	(+)	(+)	(+)	(+)	(+)	(+)
RHA	(+)	(+)	(+)	(+)	(+)	(+)	(+)
Sucrose	(+)	(+)	(+)	(+)	(+)	(+)	(+)
Melobiose	(+)	(+)	(+)	(+)	(+)	(+)	(+)
AMY	(-)	(-)	(-)	(-)	(-)	(-)	(-)
Arabinose	(+)	(+)	(+)	(+)	(+)	(+)	(+)
H. Sensibilité au tabac	(+)	(+)	(+)	(+)	(+)	(+)	(+)
Oxydase	(-)	(-)	(-)	(-)	(-)	(-)	(-)
ADH	(+)	(+)	(+)	(+)	(+)	(+)	(+)
H.L	(+)	(+)	(+)	(+)	(+)	(+)	(+)
YDC	(+)	(+)	(+)	(+)	(+)	(+)	(+)
NA	(+)	(+)	(+)	(+)	(+)	(+)	(+)
ONPG	(+)	(+)	(+)	(+)	(+)	(+)	(+)
LDC	(-)	(-)	(-)	(-)	(-)	(-)	(-)
ODC	(-)	(-)	(-)	(-)	(-)	(-)	(-)
CIT	(+)	(+)	(+)	(+)	(+)	(+)	(+)
H2S	(-)	(-)	(-)	(-)	(-)	(-)	(-)
URE	(-)	(-)	(-)	(-)	(-)	(-)	(-)
TDA	(+)	(+)	(+)	(+)	(+)	(+)	(+)
Catalase	(+)	(+)	(+)	(+)	(+)	(+)	(+)

ONPG: determination of enzyme beta-galactosidase presence, ADH: transformation of arginine by dishydrolase argnine, LDC: transformation of lysine by decarboxylase lysine, ODC: transformation of ornithine by decarboxylase ornithine, CIT: utilization of citrate as alone source of Carbone, H2S: production of hydrogen sulfate from thiosulfate, URE: liberation of ammoniac from urea by urease, TDA: formation of indole pyruvique acid from tryptophan by desaminase tryptophan, IND: formation of indole from tryptophan, VP: formation of acetone from sodium piruvate, GEL: liquefaction of gelatin, GIU: glucose, MAN: mannitol, INO: inositol, SOR: sorbitol, RHA: rhamnose, SAC: sucrose, MEL: melibiose, AMY: amygdaline, ARA: arabinose (formation of acid by utilization of carbon hydrate tests) (-): Negative; (+): Positive.

The 2276-1, 2276-2, 2276-3, 2276-4, 2276-5 and 2276-6 strains gave a positive reaction on pathogenicity test *in vitro*. This test showed that there was development of centre soft rot after six days of incubation on onion bulbs by *P. ananatis* (Carr *et al.*, 2008; Conn *et al.*, 2012). Result of DNA quantification showed that the quantity of DNA was sufficient for PCR (25 ng /  $\mu$ l). A band of 1550 bp size was observed in agarose gel (Fig. 2). BLAST-NCBI analyses of sequences proved that 2276-1, 2276-2, 2276-3, 2276-4, 2276-5 and 2276-6 were strains of *P. ananatis* (Table 2).

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As a result, representative isolates of the

causal agent of soft rot of onion bulbs in

# Figure 2 Electrophoritic profile of *P. ananatis* strains: (1) 2276-1. (2) 2276-2. (3) 2276-3. (4) 2276-4. (5) 2276-5. (6) 2276-6. (7) *P. ananatis* reference strain. (T-) Negative control (H2O).

100bp 1

1500

Table 2 The sequence identity/similarity.

The sequence of isolates strains (2276-1, 2276-2, 2276-3, 2276-4, 2276-5 and 2276-6)	Sequence identity / Similarity	GenBank Accession Numbers
GCGTGTATGAGAAGGCCTTCGGG TTGTAAAGTACTTTCAGCGGGGA GGAAGGCGATGTGGTTAATAACC GCATTGATTGACGTGGGTTAATAACC GCATTGATTGACGTTACCCGCAG AAGAAGCACCGGCTAACTCCGTG CCAGCAGCCGCGGTAATACGGAG GGTGCAAGCGTTAATCGGAATTA CTGGGCGTAAAGCGCACGCAGGC GGTCTGTTAAGTCAGATGTGAAA TCCCCGGGCTTAACTCGGAAACT GCATTTGAAACTGGCAGGCTGA GTCTCGTAGAGGGGGGGAAATT CCAGGTGTAGCGGTGAAATGCGT AGAGATCTGGAGGGGGGAAAGC GCGAAGGCGGCCCCTGGACGAA GACTGACGCTCAGGTGGAAAGC GTGGGGAGCAAACAGGATTAGAT ACCCTGGTAGTCGACGGCGCAGAGC GTGGGGAGTGGCTTCCGGAAGC GTGGGGAGTGGCTTCCGGAGGCT AACCTGGTAGTCGACGCGTAAA CGATGTCGACTTGGAGGTTGTTCC CTTGAGGAGTGGCTTCCGGAGGCT AACGCGTTAAGTCGACCGCCTGG GGAGTACGGCCGCAAGGTTAAAA CTCAAATGAATTGACGGGGGGCCC GCACAAGCGGTGGAGCATGTGGT TTAATTCGATGCAACGCGAAGAA CCTTACCTACTCTTGACATCCACA GAACTTGGCAGGAGATGCTTGGT GCCTTCGGGAGCCCTGAGACAGG TGCTGCAGGCCCTGAGACAGG TGCTGCAGGCCCTGAGACAGG TGCTGCAGGCCTGAGACCCCC GCACAAGCGTGGAGCCTGAGACAGG TGCTGCAGGCCCTGAGACAGG TGCTGCAGGCCTGAGACCGCCTG GGGTTTGGCAAGGCCTGCGCAGCACAGG TGCTGCATGGCCTGCGCAGCCC GCTTCGGCAGGCCTGCGCAGCACAGG TGCTGCATGGCCTGCGCAGCCTC GTGTTGTGAAATGTTGGGTTAAG	1 tetegggtga egagtggeg acgggtgagt aatgtetggg gatetgeceg atagagggg 61 ataacaetg gaaacggtg etaataeege ataacgtege aagaceaaag agggggaeet 121 tegggeetet eactategga tgaaceaga tgggattage tagtaggeg ggtaacggee 181 cacetaggeg acgateeta getggtetga gaggatgaee agecaeaetg gaactgagae 241 acggteeaga eteetaegg aggeageag ggggaatatt geaeaatggg geeaageet 301 atgeageeat geegetgta tgaagaagge ettegggtt taaagtaett teagegggga 361 ggaaggegat gtegttaata acegeattga ttgaegtae egecaageet gagaaggegg 421 aceetagge cageegtgta tgaagaagge ettegggtt taaagtaett teagegggga 361 ggaaggegat gtegttaata acegeattga ttgaegtae egecaagaet egegga 361 ggaaggegat gtegttaata acegeattga ttgaegtae egecaagae ggataeegget 421 aaeteegte cageageeg ggtaataegg agggtgeaag egttaacegg aattaetggg 481 egtaaagege acgeaggegg tetgttaagt cagatgtgaa ateceeggge ttaacetggg 541 aaetgeattt gaaaetggea ggettgggta cagatgggg ggtgagaatt ecaggtgga 601 eggtgaaatg egtagagate tggaggaata eeggtggega aggeegeee etgaegaag 661 actgaegete aggtgegaaa gegtggggg caaacaggat tagataeee ggaegaag 661 actgaegete aggtgegaaa gegtggggg caaacaggat tagataeee ggaegaag 661 actgaegete aggtgegaaa gegtggggg caaacaggat tagataeee ggaegaag 661 actgaegete aggtgegaat etggaggttgt eeettgagga gtggetteeg gagetaaege 721 geegtaaae g atgtegaett ggaggttgt eeettgagga gtggetteeg gagetaaege 781 gttaagtega eegeetggg agtaeggeeg caaaggtaa acteaaatga attgaeggg 841 geeegeacaa geggtggage atgtggttt aattegatgea acgegaagaa eettaeetae 901 tettgaeate cacagaaett ageaggaatg ettegtgee ttegggaaet etgagaeag 961 tgetgeatgg etgtegteag etegtgttg gaaagtggg gaagaageaggeggtaeae 1021 caaceettat eetttgtte cagegaatte gtegggaaat caaaggagaa ggeeggag 961 tgetgeatgg tageegaa aggeggga gaagtggag gagetgaag egecteataa 1201 agtgegtegt agteeggaa gaggtggga tgaegteag ecectaegga aggeegga gaacteaga 1201 agtgegga taggtggea egagtegga tgaegtega acteggege ettgtaeaa eegeegga 1221 caceatggga gtgggttgea aaagaagag gageetgege ettgtaeaa eegeegga gagetgaa 1321 caceatggga gtgggttgea aaagaagtag gtagettaae etteggagg geeettaeca 1381 etttgtgatt catgaetggg gtgaagtegt aacaaggtag gtagetga e	AY7411 61.1

### The sequences of the isolates strain amplified by RNA 16S, present a 99% similarity with accession number of AY741161.1.

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To our knowledge, this is the first report of bacterial bulbs soft rot of onion in Morocco caused by *P. ananatis*. Because the soft rot of onion is the most destructive disease during storage, this study is considered as a basis for works that will target the *P. ananatis* ecologic and behaviors, to solve the soft rot problem by applying a biological control by using antagonistic bacteria. A development of a molecular method is very important for a rapid detection of *P. ananatis* in soft rot bulbs.

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# اولین گزارش از ردیابی Pantoea ananatis، عامل پوسیدگی باکتریایی مغز پیاز در مراکش

الحسن آچبانی'، حمید مزوز'، عبدالطیف بزوآزا' و سومیا صدیق'' \*\*

۱ – آزمایشگاه گیاهپزشکی، مکنز، مراکش. ۲ – آزمایشگاه بیوتکنولوژی گیاهی و بیولوژی مولکولی، دانشکده علوم، مکنز، مراکش. \* پست الکترونیکی نویسنده مسئول مکاتبه: sadik.soumia@hotmail.com دریافت: ۳ آذر ۱۳۹۳؛ پذیرش: ۱۲ آبان ۱۳۹۴

چکیده: گونه Pantoea ananatis یک بیمارگر مهم عامل پوسیدگی مغز پیاز میباشد. استرینهای جدا شده از غدههای پیاز با علایم پوسیدگی نرم براساس ویژگیهای مورفولوژیک برای واکنش فوق حساسیت روی توتون آزمایش شدند. براساس آزمونهای بیوشیمیایی استرینهای بیماریزا P. ananatis تشخیص داده شد و با تعیین توالی ناحیه RNA 16S ریبوزومی نتایج تأیید شد. با مایهزنی این گونه روی برگها و غده پیاز علایم مشابه مشاهده شد. این بیماری اولین گزارش از وقوع باکتری P. ananatis

واژگان كليدى: Allium cepa ، پوسيدگى باكتريايى مغز، مراكش، پياز، Allium cepa