1	Virginia Ferreira <sup>1</sup> , Matías González <sup>2</sup> , María Julia Pianzzola <sup>1</sup> , Núria S. Coll <sup>3</sup> , María Inés Siri <sup>1*</sup> and Marc
2	Valls <sup>3,4*</sup>
3	
4	<sup>1</sup> Área Microbiología, DEPBIO, Facultad de Química, Universidad de la República, Uruguay.
5	<sup>2</sup> Instituto Nacional de Investigaciones Agropecuarias (INIA), Estación Experimental Salto Grande,
6	Salto, Uruguay.
7	<sup>3</sup> Centre for Research in Agricultural Genomics (CSIC-IRTA-UAB-UB), Bellaterra, Catalonia, Spain
8	<sup>4</sup> Department of Genetics, University of Barcelona, Barcelona, Catalonia, Spain
9	
10	* Corresponding authors: <u>msiri@fq.edu.uy;</u> <u>marcvalls@ub.edu</u>
11	
12 13	Running head: Detection of latent infections in potato breeding programs

# Molecular detection of *Ralstonia solanacearum* to facilitate breeding for resistance to bacterial wilt in potato

17

#### 18 Abstract

19 Potato bacterial wilt is caused by the devastating bacterial pathogen Ralstonia solanacearum. 20 Quantitative resistance to this disease has been and is currently introgressed from a number of wild 21 relatives into cultivated varieties through laborious breeding programmes. Here, we present two 22 methods that we have developed to facilitate the screening for resistance to bacterial wilt in potato. 23 The first one uses *R. solanacearum* reporter strains constitutively expressing the *luxCDABE* operon or 24 the green fluorescent protein (gfp) to follow pathogen colonisation in potato germplasm. 25 Luminescent strains are used for non-destructive live imaging, while fluorescent ones enable precise 26 pathogen visualisation inside the plant tissues through confocal microscopy. The second method is a 27 BIO-multiplex-PCR assay that is useful for sensitive and specific detection of viable R. solanacearum 28 (IIB-1) cells in latently infected potato plants. This BIO-multiplex-PCR assay can specifically detect 29 IIB-1 sequevar strains as well as strains belonging to all four R. solanacearum phylotypes and is 30 sensitive enough to detect without DNA extraction 10 bacterial cells per ml in complex samples.

31 The described methods allow the detection of latent infections in roots and stems of asymptomatic
32 plants and were shown to be efficient tools to assist potato breeding programs.

33

Key words: Bacterial wilt, potato brown rot, *Ralstonia solanacearum*, *Solanum tuberosum*, plant
breeding, disease resistance.

36

#### 37 **1.** Introduction

38 Bacteria can cause a range of diseases in economically important crops, leading to important losses.
39 *Ralstonia solanacearum*, the causal agent of bacterial wilt, also referred as brown rot in potato, is one
40 of the most devastating plant pathogens worldwide [1]. A recent taxonomic revision has led to the
41 distinction of three separate species within the species complex [2]. In cold and temperate regions of

the world, potato crops are mainly affected by *R. solanacearum* strains belonging to the phylotype
IIB, sequevar 1 [3].

Potato bacterial wilt disease control is difficult due to pathogen persistence in water, soil and latently infected symptomless tubers. The use of resistant or tolerant potato varieties combined with preventive measures throughout an integrated pest management approach is highly recommended [4]. Breeding for resistance to *R. solanacearum* in Solanaceae is challenging and must combine durable resistance with desirable agronomic traits. In potato, breeding for resistance to bacterial wilt has been successfully introgressed from *S. phureja* [5], and the highly resistant wild potato *S. commersonii* Dun is currently being used [6, 7, 8].

51 Knowledge on pathogen distribution and multiplication in plant tissues is critical to fully exploit the 52 potential of sources of bacterial wilt resistance through breeding programs. Asymptomatic latent 53 infections should also be considered to avoid the selection of tolerant varieties which promote 54 pathogen dissemination under favourable environmental conditions [9].

55 We present here two methodologies that we have developed to evaluate bacterial loads in inoculated 56 potato germplasm: the use of reporter strains [10, 11] and a BIO-multiplex-PCR assay.

57

#### 58 **2.** Materials

59

#### 2.1 Plant varieties and plant growth materials.

- The potato cultivar *S. tuberosum* cv. Chieftain is used as a susceptible control. Susceptible
  (13001.79, 13001.107) and resistant (1201.27, 09509.6) interspecific potato clones derived
  from different breeding lines are selected from the National Institute for Agricultural
  Research (INIA, Uruguay) germplasm collection.
- Potato multiplication medium (PMM): mix 20 ml of nitrate solution (95 g/L KNO<sub>3</sub>, 82,5 g/L
   NH4SO<sub>3</sub>. Adjust to 1 L with sterile distilled water), 20 ml of sulphate solution (18,5 g/L
   MgSO<sub>4</sub>·7H<sub>2</sub>O, 1,25 mg/L CuSO<sub>4</sub>·5H<sub>2</sub>O, 1,115 g/L MnSO<sub>4</sub>·4H<sub>2</sub>O, 0,43 g/L ZnSO<sub>4</sub>·7H<sub>2</sub>O. Adjust
- 67 to 1 L with sterile distilled water), 20 ml of halogens solution (22 g/L CaCl<sub>2</sub>·H<sub>2</sub>O, 41,5 mg/L

68		KI, 1,25 mg/L CoCl <sub>2</sub> ·6H <sub>2</sub> O. Adjust to 1 L with sterile distilled water), 1 ml of vitamin solution
69		(0,05 % folic acid, 0,1% biotin, 0,1% choline chloride, 0,1% pantothenic acid, 0,1% thiamine,
70		0,2 % nicotinamide, 0,2% pyridoxine, 0,2% aminobenzoic acid), 0,17 g/L KH2PO4, 6,2 mg/L
71		H3BO3, 0,25 mg/L Na2MoO4·2H2O, 37,9 mg/L Na2EDTA.2H2O, 27,8 mg/L FeSO4.7H2O, 30
72		g/L sucrose, 0,2 g/L myo-inositol and 8 g/L agar. Adjust to pH 5,8 with KOH and to 1 L with
73		sterile distilled water.
74	3.	Glass containers with autoclavable plastic lid.
75	4.	Sterile glass petri dishes.
76	5.	Metal clamps and scalpels.
77	6.	Metal sterilizer.
78	7.	Laminar flow cabinet.
79	8.	In vitro growth chambers for healthy plants at 22°C with cycles of 16 h light/8 h darkness.
80	9.	Soil mix (Tref Substrate).
81	10	. 5 cm <sup>3</sup> Multicell-trays.
82	11	. 170-cm <sup>3</sup> individual pots.
83	12	. Greenhouse with natural light and room temperature.
84	13	. Growth chambers with controlled temperature, humidity and light.
85		
86	2.2	Bacterial strains and bacterial culture.
87	1.	R. solanacearum UY031 strain [12], R. solanacearum reporter strains UY031 Pps-GFP and
88		UY031 Pps-lux [10, 11].
89	2.	Rich B medium: 10 g/L Bacteriological peptone, 1 g/L Yeast extract and 1 g/L Casamino
90		Acids. Add 1.5% Agar for solid media before autoclaving. Before plating, add 0.5% Glucose
91		and 0.005% Triphenyltetrazolium chloride (TTC). Adjust to pH 7,0. Supplement with
92		gentamicin at 5 and 75 $\mu\text{g/ml}$ in liquid and solid cultures respectively for selection of
93		reporter strains.
94	3.	mSMSA medium [13]: 10 g/L Bacteriological peptone, 5 ml of glycerol and 1 g/L Casamino

95	Acids. Add 1.5% Agar for solid media before autoclaving. Before plating, add 10 ml of
96	polymyxin B sulfate (1%), 10 ml of cyclohexide (1%), 2,5 ml of bacitracin A (1%), 500 $\mu$ l of
97	penicillin (0,1%), 500 $\mu$ l of cloranfenicol (1%), 500 $\mu$ l of crystal violet (1%) and 0.005% TTC.
98	Adjust to pH 7,0.
99	4. Sterile Petri dishes.
100	5. Sterile tubes.
101	6. Sterile saline solution: add 9 g/L NaCl in distilled water before autoclaving.
102	7. Spectrophotometer.
103	
104	2.3 Evaluation of bacterial colonisation using reporter strains.
105	1. Luminometer (LAS4000 light imager system, FujiFilm).
106	2. Epifluorescence microscope.
107	3. Confocal fluorescence microscope.
108	4. Glass slides.
109	5. Coverslips.
110	6. Solid vaseline.
111	7. 20 ml syringe.
112	8. 200 μl pipette tips.
113	9. Razor blade.
114	10. Agarose 1 %.
115	
116	2.4 BIO-multiplex-PCR.
117	1. Ethanol 70%.
118	2. Sterile water.
119	3. Sterile absorbent paper.
120	4. Sterile plastic bags.
121	5. 50mM phosphate buffer: 4,26 g/L Na2HPO4 and 2,72 g/L. Adjust to pH 7,0 and to 1 L with

122	sterile distilled water.	
123	6. Laboratory blender.	
124	7. 1000 and 200 μl pipettes with sterile tips.	
125	8. Sterile glass beads for liquid dissemination on Petri dishes.	
126	9. 28°C Incubator.	
127	10. Sterile loops.	
128	11. Sterile tubes.	
129	12. Centrifuge.	
130	13. PCR tubes.	
131	14. Thermocycler.	
132	15. 5 units/ $\mu$ l Taq DNA polimerase and its concentrated buffer.	
133	16. 25 mM MgCl <sub>2</sub>	
134	17. 5 mM Deoxyribonucleotide triphosphates (dNTPs).	
135	18. 10 $\mu$ M of IIB-1 specific primers (00876F: 5'-GGATTCAAGGTATCGC	CAGA-3'; 00876R: 5'-
136	CATAGCCGCTTCTTCTTGG-3') and general primers	[14] (759: 5′-
137	GTCGCCGTCAACTCACTTTCC-3'; 760: 5'-GTCGCCGTCAGCAATG	CGGAATCG-3′).
138	19. MiliQ water.	
139		
140	3. Methods	
141	3.1 Plant growth	
142	1. Start <i>in vitro</i> cultures from <i>in vitro</i> plants containing 3-4 internodes g	rown <i>in vitro</i> in tubes
143	or glass bottles with PMM medium.	
144	2. Cut single-node pieces of plants and transfer them into glass both	tles with fresh PMM
145	previously autoclaved (see Note 1). Put around 10-12 single-node piece	es per glass bottle (Fig.
146	1a).	
147	3. Leave plantlets growing for 15-20 days in the in vitro growth chamber	at 22°C with cycles of

- 148 16 h light/8 h darkness.
- 4. Before transfer to soil, incubate closed glass bottles with plants in a greenhouse under natural
  light and room temperature during 3-5 days for acclimatisation.
- 151 5. Transfer plants to soil either in plastic trays with 5-cm<sup>3</sup> wells for bacterial wilt resistance
  152 screening (Fig. 1b), or to individual pots with 170-cm<sup>3</sup> for evaluation of bacterial colonisation
- by inoculation with reporter strains (Fig. 1c) (*see* **Note 2**).
- 154 6. Incubate for 15-20 days in the greenhouse under natural light and room temperature, until
  155 plants reach a height of 10 cm.
- 156 7. Water plants frequently checking that soil mix is wet but avoiding flooding.
- 157 8. Before inoculation, incubate plants 2-3 days for acclimation in the growth chamber at 28 °C
- 158 with cycles of 16 h light/8 h darkness and 65 % of humidity.

- 160 **3.2** Bacterial culture and inoculum preparation
- Streak bacterial strains from glycerol stocks kept at -80 °C on B medium. Grow for 2 days at
   28 °C. Supplement the medium with gentamicin when reporter strains are used.
- 163 2. Pick a single colony to inoculate 20 ml of liquid rich B medium and incubate overnight at 28
- 164 °C with orbital shaking (200 rpm) (*see* **Note 3**).
- 165
  3. Centrifuge the bacterial culture, discard the supernatant and carefully re-suspend the pellet
  166 with 20 ml of sterile saline solution or sterile water.
- 167 4. Using a spectrophotometer, measure the optical density at 600 nm (OD<sub>600</sub>) of the cell
  168 suspension.
- Add the required volume of cell suspension for adjusting the desired volume of inoculum to
  a final concentration of 10<sup>7</sup> cfu/ml (consider that an OD<sub>600</sub> of 0.1 corresponds to 10<sup>8</sup> cfu/ml).
- 1716. Confirm the final inoculum concentration by preparing 10-fold dilutions and colony172counting in rich B medium plates.

#### 174 **3.3** Evaluation of bacterial colonisation using reporter strains

- Two types of *R. solanacearum* reporter strains are used: a bioluminescent strain for nondisruptive, macroscopic assessment of bacterial colonisation, and a fluorescent strain for
- 177 microscopic evaluation of colonisation at the tissue level.
- 178 **3.3.1** Evaluation of colonisation using a luminescent strain.
- For evaluation of bacterial colonisation, use 5-10 replicate plants for each clone, grown in
   individual pots and arranged using a completely randomized design (Fig. 1c).
- Prior to inoculation, damage roots slightly by making three holes of 2 cm deep in the soil of
   each pot with a disposable 1000 μl pipette tip.
- 183
  3. Inoculate potato clones by drenching, using 40 ml of bacterial suspension of *R. solanacearum*184
  UY031 Pps-lux strain to reach a final density of 10<sup>6</sup> cfu/g of soil. Inoculate plants with saline
- 184 UY031 Pps-lux strain to reach a final density of 10<sup>6</sup> cfu/g of soil. Inoculate plants with saline
  185 solution as negative control treatment.
- 4. After inoculation, record daily, for 6-10 days, bacterial colonisation of plant tissues using the
  Fuji Film LAS4000 light imager system with the chemiluminescence settings of incremental
- 188 exposure time each 2 min and sensitivity/resolution set to high binning (Fig. 2).
- 189

#### 190 **3.3.2** Evaluation of colonisation using a fluorescent strain.

- Use the same plant inoculation procedure described for the luminescent strain using instead
   *R. solanacearum* UY031 Pps-GFP strain.
- Collect root and stem samples 2- and 7-days post inoculation (dpi) to follow pathogen
   colonisation.
- 195 3. Remove plants from pots and wash roots with tap water to remove adherent soil.
- 4. Surface sterilize stem and roots of each plant with ethanol 70% for 1 min, rinse with sterile
  water for 1 min and dry with sterile paper towels in a laminar flow cabinet.
- 198 5. Select colonized roots to be observed by confocal microscopy and put them on a glass slide
- 199 with agarose 1% to maintain sample hydrated during manipulations (*see* **Note 4**).

200	6.	Put solid vaseline in the border of a coverslip with syringe, making a 2-3 mm wide retaining
201		wall (see Note 5).
202	7.	Place the selected colonized roots (see step 5) on the coverslip surrounded by solid vaseline.
203	8.	Cut 2-cm stem segments using a previously disinfected scalpel, cut from 1 cm above root.
204		Make six to 10 cross-sections by hand with razor blade on the end of each stem segment and
205		place them on the coverslip with root segments of each plant.
206	9.	Add melted agarose 1% as immersion medium surrounding stem and root segments. Vaseline
207		wall should retain melted agarose (see Note 6).
208	10.	Immediately place a glass slide to seal the chamber (Fig. 3a).
209	11.	Observe stem cross-sections and roots using a confocal ( <i>see</i> <b>Note 7</b> ) microscope (Fig. 3b).
210		
211	3.4	Bacterial wilt resistance evaluation.
212	1.	Use two replicate trays, each containing eight plants per clone. Arrange the trays in the
213		growth chamber using a completely randomized design.
214	2.	Prior to inoculation, damage roots slightly by making one hole of 2 cm deep in the soil of
215		each well with a disposable 1000 $\mu$ l pipette tip.
216	3.	Inoculate each cell by adding 1 ml of bacterial suspension adjusted to 10 <sup>7</sup> cfu/ml. Inoculate
217		plants with saline solution as negative control treatment.
218	4.	Record wilting symptoms until almost all the susceptible control plants (cv. Chieftain) are
219		totally wilted (approximately 28 dpi). Disease scoring is performed using a semi-quantitative
220		scale that ranges from 0 to 4, in which $0=$ no wilting, $1 = 25\%$ of the leaves wilted, $2=50\%$ of
221		the leaves wilted, $3=75\%$ of the leaves wilted, $4=100\%$ of the leaves wilted, dead plant ( <i>see</i>
222		Note 8).
223	5.	Estimate the resistance level by calculating the area under disease progression curve
224		(AUDPC) based on the average wilt scoring for each clone.
225		
226	3.5	Evaluation of latent infections using BIO-multiplex-PCR.

- The occurrence of latent infections is determined in genotypes with 0-30% of wilted plants
   after 28 dpi (evaluated using Method 3.4).
- 2. Collect only asymptomatic plants and wash with tap water to remove the adherent soil.
  230 Surface sterilise the plants with ethanol 70% for 1 min, rinse with sterile water for 1 min and
  231 dry with sterile absorbent paper towels.
- 3. Using a sterile scalpel, cut 2 cm stem segments from the basal part of each plant, just above
  the soil level.
- 4. Pool stem segments from each replicate tray of each genotype and place them into a sterile
  bag. Weigh stem pools, add phosphate buffer (10 ml per gram of tissue) and mix in a
  laboratory blender at high speed for 10 min.
- 237 5. Spread aliquots of 100 μl of stem extract onto each of two plates of mSMSA and incubate at
  238 28 °C.
- After 48 h wash one of the mSMSA plates with 2 mL of sterile water (*see Note 9*). The
  remaining plate is maintained at 28 °C for visual recovery and enumeration of *R. solanacearum* colonies.
- Centrifuge washed suspensions at 8000 rpm for 5 min and re-suspend pellets in 100 µl of
  sterile water in Eppendorf tubes. Boil for 20 min and store on ice until used as template for
  amplification by multiplex-PCR.
- 8. Multiplex PCR: mix 5 μl of lysate; 1,5 mM MgCl2; 0,2 mM each of four dNTPs; 10 pmol of
  each IIB1-specific primer (00876F/00876R); 10 pmol of primers 759/760; 1,5 U Taq DNA
  polymerase and the buffer supplied with the enzyme in 25 μl of reaction volume.
- Amplification program: 5 min at 96 °C, followed by 40 cycles of 94 °C for 30 s, 60 °C for 30 s,
  and 72 °C for 45 s, with a final extension step for 10 min at 72 °C.
- 250 10. See results by gel electrophoresis using agarose 1.5% for detection of both amplification
  251 products. IIB-1 specific primers 00876F/00876R and 759/760 amplify products of 342 bp and
  252 280 bp, respectively (Fig. 4).

- 253 11. Record the number of *R. solanacearum* colonies grown in the remaining mSMSA plate after
  254 7-10 days of incubation at 28 °C. Latently infected plants usually have a pathogen
  255 concentration of 10<sup>3</sup> to 10<sup>6</sup> cfu/g of stem tissue.
- 256
- 257 **4. Notes.**
- Carry out micro-propagation procedures inside the laminar flow cabinet. Cut single-node
   pieces with metal clamps and scalpels inside glass petri dishes previously autoclaved. Sterilize
   metal clamps into the metal sterilizer every 3-4 plants. Before autoclaving, melt PMM and
   transfer 30-50 ml inside the glass bottles. Single-node stem pieces should be in contact with
   PMM for plant growth.
- 263
  2. Before transfer plants to soil, remove agar from roots carefully avoiding root damage. Rinse
  264
  plants in tap water 2-3 times until no agar remains adhere to the roots.
- 265 3. Pick a colony with the typical morphology of *R. solanacearum*: large, fluidal, and either
  266 entirely white or white with a red centre.
- 267
  4. Put whole root system on a glass slide with agarose 1% to observe roots on epi-fluorescence
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  269
- 270 5. Put a 200 μl pipette tip in the syringe tip to make a finest solid vaseline retaining wall.
- Add agarose immediately after placing stem cross-sections. Tissue segments could dry out if
  agarose addition is delayed.
- Fluorescence can also be detected using an epifluorescent microscope, although, interference
  with the chlorophyll autofluorescence often makes interpretation difficult. The intensity of
  the GFP signal and the microscope fluorescence filters are key variables that have to be
  evaluated in each case. The use of a confocal microscope highly improves signal specificity
  and resolution and is always effective.

278		8. Scoring of disease symptoms has some degree of subjectivity. It is strongly recommended
279		that the same person carries out the whole symptom recording to avoid experimental bias.
280		9. The mSMSA plate should be washed only when no evident growth or pin-point colonies are
281		observed after 48 hs of incubation. If confluent growth is observed the washing should be
282		replaced by collecting the cells with a sterile toothpick and diluting in 100 $\mu l$ of sterile water,
283		before following with the lysis and amplification procedure by multiplex PCR.
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292		nov., R. solanacearum phylotype IV strains as Ralstonia syzygii subsp. indonesiensis subsp. nov.,
293		banana blood disease bacterium strains as <i>Ralstonia syzygii</i> subsp. <i>celebesensis</i> subsp. nov. and <i>R.</i>
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   321 strain-specific DNA probes and PCR primers for identifying *Burkholderia solanacearum* 322 (formerly *Pseudomonas solanacearum*). Asia Pac J Mol Biol Biotechnol 5:19-30.
- 323

#### 324 Figure legends

- 325 Fig 1. Representative photographs of the different potato plant growth stages. (a) In vitro plants
- 326 growing on PMM in glass bottles. (b) Plants grown in plastic multicell-trays for bacterial wilt
- 327 resistance evaluation. (c) Plants grown in individual pots for inoculation with reporter strains.

Fig 2. Bioluminescence imaging of *Ralstonia solanacearum* UY031 Pps-lux strain coloninising
different potato genotypes. (a) Susceptible potato cultivar *Solanum tuberosum* cv. Chieftain. (b)
Resistant potato clone 11201.27. Light gray indicates background luminescence and black regions
are colonized tissue by light-emitting bacteria. Images were acquired at 3 and 6 days post
inoculation (dpi).

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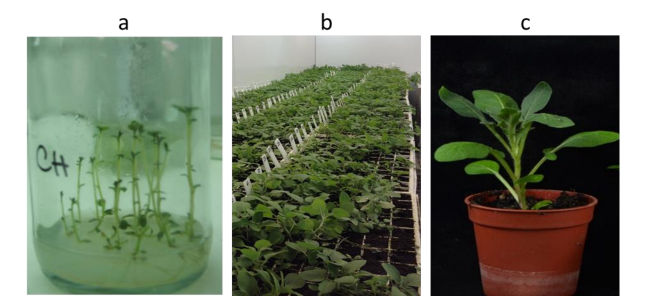
Fig 3. Evaluation of bacterial colonisation using the *Ralstonia solanacearum* fluorescent reporter strain UY031 Pps-gfp. (a) Confocal visualisation chamber containing stem cross-sections and root segments. (b) Representative confocal fluorescence micrographs of stem cross-section (Resistant potato clone 09509.6, left) and root (Susceptible potato clone 13001.79, right). Images were acquired 7 days post inoculation (dpi). Dark arrows show bacterial colonisation.

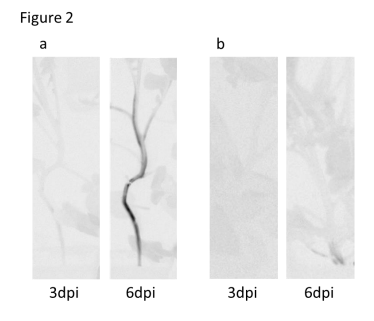
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Fig 4. Gel electrophoresis showing positive results of the multiplex-PCR for detection of latent
 infections in asymptomatic potato plants inoculated with *Ralstonia solanacearum* (IIB-1).

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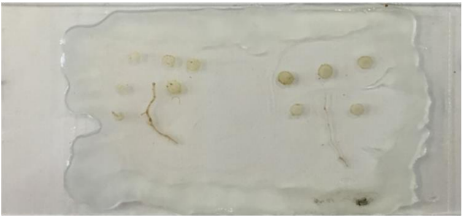
### Figure 1





## Figure 3

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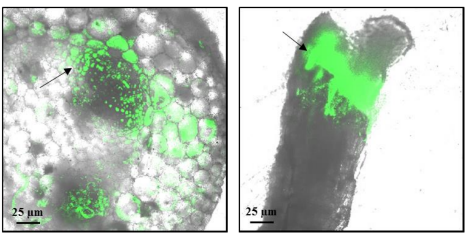


Figure 4

