# RESEARCH

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# Root exudate profiling of maize seedlings inoculated with *Herbaspirillum seropedicae* and humic acids

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# Abstract

**Background:** Co-inoculation of maize with *Herbaspirillum seropedicae* and humic substances increases the sizes of plant-associated bacterial populations and enhances grain yields under laboratory and field conditions. Root exudation is a key mechanism in the regulation of plant-bacterial interactions in the rhizosphere; humic matter supplementation is known to change the exudation of H<sup>+</sup> ions and organic acids from maize roots. Our starting premise was that *H. seropedicae* and humic acids would modify maize seedling exudation profiles. We postulated that a better understanding of these shifts in exudate profiles might be useful in improving the chemical environment to promote better performance of plant growth-promoting bacteria delivered as bioinoculants. Thus, root exudates of maize were collected and analyzed by gas chromatography-mass spectrometry (GC-MS) and proton nuclear magnetic resonance (<sup>1</sup>H NMR).

**Results:** Nitrogenous compounds, fatty acids, organic acids, steroids, and terpenoid derivatives were the main structural moieties found in root exudates. Significant changes in exudation patterns occurred 14 days after the initiation of experiments. Quantities of fatty acids, phenols, and organic acids exuded by seedlings treated with humic acids alone differed from the quantities exuded in other treatments. Seedlings treated with *H. seropedicae* or *H. seropedicae* in combination with humic acids exuded a diversity of nitrogenous compounds, most of which had heterocyclic structures. Twenty-one days after initiating the experiment, seedlings treated with *H. seropedicae* alone exuded elevated quantities of steroids and terpenoid derivatives related to precursors of gibberellic acids (kaurenoic acids).

**Conclusions:** Changes in root exudation profiles induced by our treatments became most marked 14 and 21 days after initiation of the experiment; on those days, we observed (i) increased fatty acid exudation from seedlings treated only with humic acids and (ii) increased exudations of nitrogenated compounds and terpenes from seedlings treated only with *H. seropedicae*. Improved knowledge on the effects of bacterial inoculants and supplementation with humates on plant exudate composition may contribute substantially to improved understanding of plant metabolic responses and lead to new approaches in the use of selected compounds as additives in bioinoculant formulations that will modulate the cross-talk between bacteria and plants, thereby improving crop yields.

Keywords: Plant growth-promoting bacteria; Endophytic interaction; Humic substances

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# Background

Root exudates function in processes of plant adaptation. They have roles in nutrient cycling in the rhizosphere and in responses to pathogens and symbiotic microorganisms [1]. The quantities of organic compounds exuded by roots are variable, but they are frequently a significant proportion of the carbon fixed photosynthetically by plants [2]. Plant type, species, age, and environmental factors, including biotic and abiotic stressors, all affect exudation profiles [3,4]. The plant rhizosphere modulates microbial community structure and function, primarily through the release of chemical compounds [5].

The dominant organic compounds exuded by roots reflect central components of cell metabolism, including free sugars (e.g., glucose, sucrose), amino acids (e.g., glycine, glutamate), and organic acids (e.g., citrate, malate, and oxalate) [2]. Maize exudates comprise sugars (70%), phenolics (18%), organic acids (7%), and amino acids (3%) [6]. Other compounds, like fatty acids, sterols, enzymes, vitamins, and plant growth regulators (e.g., auxins, gibberellins, and cytokinins), are generally released in only very small quantities [7]. Although only small quantities of secondary metabolites are exuded by roots, their plant role in signaling to the microbial community is substantial [8].

Recently, we determined that the combined inoculation of maize with the endophytic diazotrophic bacterium *Herbaspirillum seropedicae* and humic acids increased root colonization and promoted plant growth and grain yields [9]. Canellas and Olivares [10] reviewed the use of humic substances as plant growth promoters; they considered the effects of these substances on plant metabolism and their use as carriers in procedures for the bioinoculation of plant growth-promoting bacteria in field crop systems.

Canellas et al. [11] reported changes in maize exudation profiles after humic acid application that included enhanced secretion of inorganic ions (i.e.,  $H^+$  ions) and short-chain organic acids [12]. The addition of humic acids of different size fractions may have substantial effects on the quantities of bioavailable carbon deposited by maize plant roots; these additions produce significant changes in the structure and activity of soil microbial communities [13]. Hence, chemical changes induced by humic matter augmentation in the rhizosphere may enhance colonization of maize plants by inoculated endophytic diazotrophic bacteria carried in the humic substances.

*H. seropedicae* is a plant growth-promoting diazotrophic  $\beta$ -proteobacterium found mainly in association with grasses and other non-leguminous plants [14]. Roesch et al. [15] used molecular tools to assess diazotrophic bacterial diversity within rhizosphere soils, roots, and stems of field-grown maize and observed a predominance of  $\alpha$ -proteobacteria and  $\beta$ -proteobacteria sequences in the rhizosphere soil and stem samples; *Herbaspirillum*  was one of the dominant genera in the interiors of maize plants but was rarer in soil. The members of this genus have been tested in the formulation of biofertilizers, with variable success in field crop trials ([16-19] and references therein [20-23]). The whole genome sequence of *H. seropedicae* has been published [24]. The species' capacity for N fixation, production of auxin and other phytohormones, and the colonization of diverse plant species has been previously demonstrated [25-27].

Root colonization is a basic first step for successful inoculation. An expansion of studies on metabolite exchange between plants and bacteria and the genetic responses of plants will fill knowledge gaps in our understanding of the colonization process. The role of root-exuded flavonoids in legume-Rhizobium interactions has been examined in detail. These exudations generate a finely tuned cross-molecular dialogue involving the secretion of lipochitooligosaccharides and the modulation of bacterial cell wall surface polysaccharides (extracellular polysaccharide (EPS) and lipopolysaccharide (LPS)) that result in plant root nodulation [28]. There is less information available on the role of plant metabolites in successful interactions with non-nodulating plant growth-promoting bacteria. Gough et al. [29] showed that flavonoids promote endophytic colonization of Arabidopsis thaliana (L.) Heynh. roots by H. seropedicae, and Tadra-Sfeir et al. [30] demonstrated that naringenin (a flavonoid in the flavonone class) is involved in the gene expression of cell wall components (EPS, LPS) and auxins. In addition to its role in the genetic modulation of cell wall assembly, H. seropedicae has an operon associated with the degradation of aromatic compounds [31]. An ability to degrade flavonoids would likely confer an important competitive advantage in rhizosphere/root colonization of the host plant by providing both a carbon source and associated detoxification mechanisms. Balsanelli et al. [32] showed that surface lipopolysaccharides produced by H. seropedicae strain Smr1 are required for attachment and endophytic colonization of maize plants. They [32] found that the H. seropedicae attachment process is partially mediated by a root lectin that specifically binds N-acetyl glucosamine residues.

Recently, Marks et al. [33] demonstrated the potential of using bacterial metabolites to enhance the performance of biofertilizers, thereby opening the possibility of chemical manipulation of carriers to benefit bacterial delivery to field crops. Furthermore, metabolites exuded by host plants may help in guiding genomic studies of plant-bacterial interactions. In the present work, we examined main changes in exudation profiles of maize seedling roots induced under laboratory conditions by (i) single applications of humic acids or (ii) *H. seropedicae* or (iii) combinations of the bacteria and humic acids.

#### Methods

# Humic substances

Humic-like substances were extracted as described previously [11]. In brief, ten volumes of 0.5 mol  $L^{-1}$  M NaOH were mixed with one volume of earthworm compost under a N<sub>2</sub> atmosphere. After 12 h, the suspension was centrifuged at 5,000  $\times$  g; humic acids (HA) were extracted thrice in this manner, and the final HA pellet was de-ashed by combining it with ten volumes of a diluted mixture of HF-HCl solution (5 mL L<sup>-1</sup> HCl [12 M] + 5 mL L<sup>-1</sup> HF [48%,  $\nu/\nu$ ]). After centrifugation (5,000 × g) for 15 min, the sample was repeatedly washed with water until a negative test against AgNO<sub>3</sub> was obtained. Subsequently, the sample was dialyzed against deionized water using a 1,000-Da cutoff membrane (Thomas Scientific, Swedesboro, NJ, USA). The dialyzate was lyophilized. We then prepared a HA solution by solubilizing HA powder in 1 mL of 0.1 M mol L<sup>-1</sup> NaOH, followed by pH adjustment to 6.5 with 0.1 M HCl.

#### Microorganism used

H. seropedicae strain HRC 54 was originally isolated from sugarcane roots [34]. It has been used as part of the sugarcane inoculant developed by Embrapa (Brazilian Enterprise for Agricultural Research). The pre-inoculum was obtained after growth in DYGS liquid medium [35] for 24 h at 30°C on an orbital shaker rotating at 150 rpm. Subsequently, 20 µL of the suspension was transferred to JNFb liquid medium supplemented with NH<sub>4</sub>Cl (1 g  $L^{-1}$ ) and then grown for 36 h at 34°C on an orbital shaker rotating at 150 rpm. Cells were pelleted by centrifugation (4,000×g for 15 min) and resuspended in sterilized water at cell densities of  $10^8$  colony-forming units (cfu) mL<sup>-1</sup>. The inoculant was prepared by diluting 200 mL of bacterial suspension in 800 mL of humic acid solution at pH 6.5 to produce a final humic acid concentration of 50 mg C  $L^{-1}$ and a final bacterial concentration of  $2 \times 10^7$  cells mL<sup>-1</sup>.

The composition of JNFb medium (per liter) was as follows: malic acid (5.0 g), K<sub>2</sub>HPO<sub>4</sub> (0.6 g), KH<sub>2</sub>PO<sub>4</sub> (1.8 g), MgSO4."7H2O (0.2 g), NaCl (0.1 g), CaCl2 (0.02 g), 0.5% bromothymol blue in 0.2 N KOH (2 mL), vitamin solution (1 mL), micronutrient solution (2 mL), 1.64% Fe<sup>--</sup>EDTA so-lution (4 mL), and KOH (4.5 g). One-hundred milliliters of vitamin solution contained 10 mg of biotin and 20 mg of pyridoxol-HCl. The micronutrient solution contained (per liter) the following: CuSO<sub>4</sub> (0.4 g), ZnSO<sub>4</sub><sup>...</sup>7H<sub>2</sub>O (0.12 g), H<sub>3</sub>BO<sub>3</sub> (1.4 g), Na<sub>2</sub>MoO<sub>4</sub><sup>--</sup>2H<sub>2</sub>O (1.0 g), and MnSO<sub>4</sub><sup>--</sup>H<sub>2</sub>O (1.5 g); pH was adjusted to 5.8. For bacterial counts, we used the same media with a semisolid consistency obtained by adding 1.9 g  $L^{-1}$  of agar [14]. The bacterial population was determined by the most probable number technique (MPN); positive growth was recognized by the formation of a thick, white pellicle, replication was threefold, and density was expressed as the log of cell number  $g^{-1}$  root fresh mass after growth on JNFb N-free semisolid medium (following Döbereiner et al. [35]). The presence of *H. seropedicae* was confirmed by collecting a piece of pellicle with a platinum loop, mounting it on a slide under a coverslip, and making observations under phase contrast microscopy to determine cell shape and movement and colony appearance in JNFb solid medium, as described by Döbereiner et al. [35]. When cell shape in the pellicle material differed from that of *H. seropedicae*, we identified the microbes as native bacteria associated with maize roots.

### **Experimental**

#### Treatment of plants

Maize seeds (Zea mays L. var. Dekalb 7815) were surfacesterilized by soaking in 0.5% NaClO for 30 min, followed by rinsing and then soaking in water for 6 h. Afterward, the seeds were sown on wet filter paper and germinated in the dark at 28°C. Four days after germination, 30 maize seedlings with root length approximately 0.5 cm were transferred into 2.2-L vessels previously filled with 2 L of one-fourth-strength Furlani nutrient solution (containing 3.527 µM Ca, 2.310 µM K, 855 µM Mg, 45 µM P, 587 µM S, 25 µM B, 77 Fe, 9.1 µM Mn, 0.63 µM Cu, 0.83 µM Mo, 2.29 µM Zn, 1.74 µM Na, and 75 µM EDTA) with inorganic N content adjusted to a low concentration (100 µmol  $L^{-1}$  [NO<sub>3</sub><sup>-</sup> + NH<sub>4</sub><sup>+</sup>]). These low levels of N and P were used to simulate the low availability in highly weathered tropical soils and to avoid the inhibition of the diazotrophic bacteria. The seedlings were fixed into a perforated Teflon support with holes of 15-mm diameter in which seeds have been fitted. The system was continuously aerated by a low flux pump normally used in aquarium systems. Four treatments (n = 3 pots per treatment) were prepared by supplementing the nutrient solution with the following: 1. HA (50 mg C  $L^{-1}$ ), 2. plant growth-promoting bacteria (PGPB) H. seropedicae strain HRC 54 (final bacterial suspension of  $2 \times 10^7$  cells mL<sup>-1</sup>), 3. humic acids plus *H. seropedicae* (HA + PGPB), and 4. control (C), without any additions. Seedlings were collected 7, 14, and 21 days after inoculation. After 1 week, and each week thereafter, one half of the nutrient solution in each pot was replaced with fresh nutrient solution through the end of the experiment. The experiment was repeated thrice independently.

#### Exudate collection

Maize seedlings were removed from the pots; their roots were immersed in glass tubes filled with 50 mL of 0.01 mol  $L^{-1}$  KOH for 5 min to remove organic anions adhering to the root surfaces. We then thoroughly washed the roots with tap water followed by a final rinse in distilled water. Complete root systems of seedlings from a single pot were inserted in a glass tube (6.5-cm inner diameter (i.d.) × 15-cm tall) filled with 80 mL of ultrapure water in which to collect the root exudates.

After 2 h, we collected the suspensions containing root exudates and filtered them through 0.22-µm filter membranes to remove root detritus and microbial cells. The filtered samples were kept frozen until we concentrated them by liquid chromatography using 10 cm of reverse phase (RP) C18 LiChroprep® RP-18 (15 to 25 µm; Merck Millipore, Billerica, MA, USA) as the stationary phase in an open glass column (2.5-cm i.d. × 20-cm tall). The aqueous suspension of exudate was forced through the column under low pressure provided by an aquarium pump. Compounds were eluted from the column with methanol under gravity, and the solvent was removed under low temperature (4°C) under vacuum (Rocket Evaporator System, Genevac, Stone Ridge, NY, USA). We drove our exudate capture to exclude sugars and amino acids and collected mainly products of the secondary metabolism using the RP C18 column.

#### NMR sample preparation and data collection

For nuclear magnetic resonance (NMR) analysis, we dissolved exudate extracts in DMSO-d6 (700 µL). All spectra were recorded at room temperature on a Bruker Avance DRX 500 spectrometer equipped with a 5-mm inverse detection probe (Bruker GmbH, Rheinstetten, Germany) operating at 500.13 MHz for 1 h. For each sample, we recorded 360 scans (FIDs) with the following parameter settings: 64 k data points, pulse width 8.5 µs (90°), spectral width of 4,401 Hz, acquisition time of 7.4 s, and a relaxation delay of 1.0 s. For spectrum processing, we used 64 k points and applied an exponential multiplication associated with a line broadening of 0.3 Hz. Spectra were referenced to tetramethylsilane (TMS) at 0.0 ppm. To obtain exudate profiles for each of the treatments in the study, we pooled extracts from treatment replicates and dissolved the dried methanolic extracts in 700 µL of DMSO-d6. Dissolved extracts were transferred to a 5-mm NMR tube for analysis.

# Principal component analysis

The proton nuclear magnetic resonance (<sup>1</sup>H NMR) spectra were reduced to ASCII files using OACD software; the resulting data matrix was imported into The Unscrambler 10.1 software (www.camo.com). Signals corresponding to the solvent, TMS, and noise from water suppression were removed from the data set prior to statistical analysis. Principal component analysis (PCA) was performed by auto scaling the variables using normalization and calculation of the first derivative as a transformation procedure.

# Gas chromatography-mass spectrometry

After NMR analysis, we analyzed the exudates by gas chromatography-mass spectrometry (GC-MS). GC separations were performed on a GCMS QP2010 Plus instrument (Shimadzu, Tokyo, Japan) equipped with an Rtx-5MS WCOT capillary column (30 m × 0.25 mm; film thickness, 0.25 µm) (Restek, Bellefonte, PA, USA). The exudates were derivatized by refluxing 0.30 mg of sample for 1 h at 70°C with an excess of MeOH and acetyl chloride, dried under a stream of N<sub>2</sub>, followed by silvlation with 100 µL of N,N-bis[trimethylsilyl]trifluoroacetamide/1% trimethylchlorosilane (Superchrom, Milan, Italy) in closed vials at 60°C for 30 min. Chromatographic separation was achieved under the following temperature regimen: 60°C for 1 min (isothermal), rising by 7°C min<sup>-1</sup> to 100°C, and then by 4°C min<sup>-1</sup> to 320°C, followed by 10 min at 320°C (isothermal). Helium was the carrier gas supplied at 1.90 mL min<sup>-1</sup>, the injector temperature was 250°C, and the split injection mode had a split flow of 30 mL min<sup>-1</sup>. Mass spectra were obtained in EI mode (70 eV) scanning in the range of m/z 45 to 850 with a cycle time of 1 s. Compound identification was based on comparisons of mass spectra with the NIST library database (http://www.nist.gov/srd/ nist1a.cfm), published spectra, and real standards. Due to the large variety of compounds with different chromatographic responses that we detected, external calibration curves for quantitative analysis were built by mixing methyl esters and/or methyl ethers of the following molecular standards: tridecanoic acid, octadecanol, 16-hydroxyhexadecanoic acid, docosandioic acid, β-sitosterol, and cinnamic acid. Increasing quantities of standard mixtures were loaded into a quartz boat and moistened with 0.5 mL of tetramethylammonium hydroxide (TMAH) solution (25% in methanol).

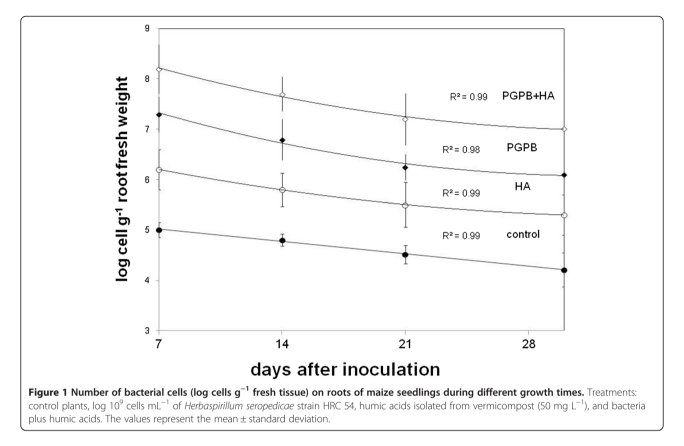
# Results

# Root tissue colonization by H. seropedicae strain HRC54

We examined the population dynamics of *H. seropedicae* strain HRC 54 associated with maize roots 7, 14, 21, and 30 days after inoculation (Figure 1). For all inoculation treatments (PGPB and HA + PGPB), the root-associated bacterial numbers were higher than those of uninoculated plants (controls). Cell shape and colony appearance confirmed the presence of *Herbaspirillum* in the pellicle harvested from the highest dilution, thereby indicating the effectiveness of inoculation.

Maize plants treated with only HA had higher bacterial numbers associated with roots than control plants. Even after seed surface disinfection, diazotrophic bacteria were recoverable from treated plants; these microbial populations were naturally occurring N fixers associated with maize seeds. They were clearly different from *H. seropedicae* (under phase contrast microscopy) in cell shape and colony form when grown in JNFb solid medium (data not shown).

We compared treatments PGPB and HA + PGPB, observing higher numbers of root-associated viable *H. seropedicae* cells in the latter. This result is qualitatively similar



to those obtained previously [9], indicating that HA *per se* increase the numbers of *H. seropedicae* cells colonizing root tissues and help maintain large populations in inoculated plants over protracted time periods (Figure 1). We identified similar tendencies for natural diazotrophs, whose population sizes were enhanced by HA application (in comparison with populations in control plants).

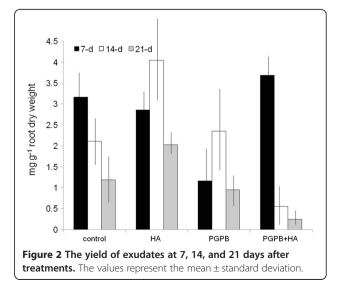
#### Mass of root exudates

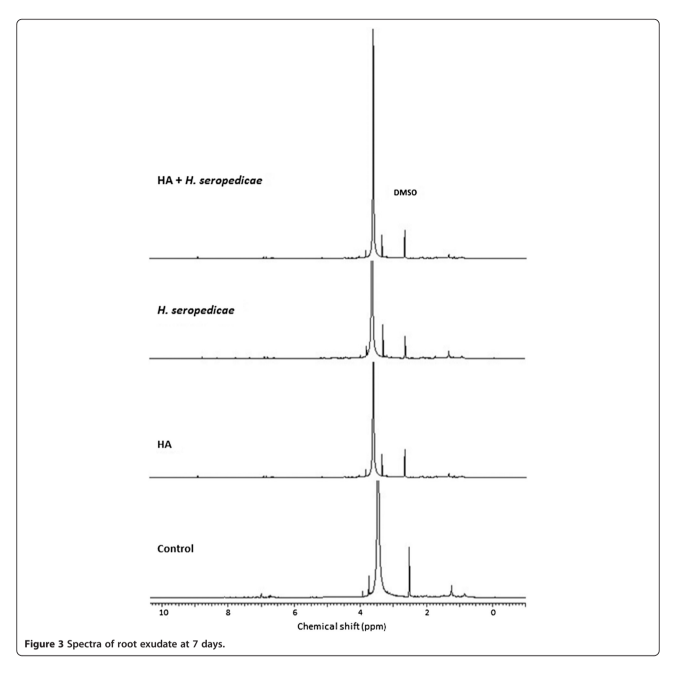
The yields of exudates retained by the procedures used are depicted in Figure 2. Over the course of the experiment, the quantities of exudates collected across sampling occasions and treatments ranged broadly between 0.25 and 4.00 mg g<sup>-1</sup> root dry weight, making it difficult to identify any treatment effects.

# <sup>1</sup>H NMR and PCA

Nuclear magnetic resonance (<sup>1</sup>H NMR) coupled with multivariate analysis (PCA) enabled rapid discrimination among root exudate samples. The <sup>1</sup>H NMR spectra of the exudate compounds eluted from the RP C18 column with methanol on days 7, 14, and 21 are depicted in Figures 3,4,5, respectively; the main <sup>1</sup>H chemical shifts are detailed in Table 1. Visual inspection of the <sup>1</sup>H NMR spectra revealed a predominance of signals in the carbohydrate region (2.5 to 4.5 ppm) followed in rank order by signals in the aliphatic/organic acid (0.0 to 3.0 ppm)

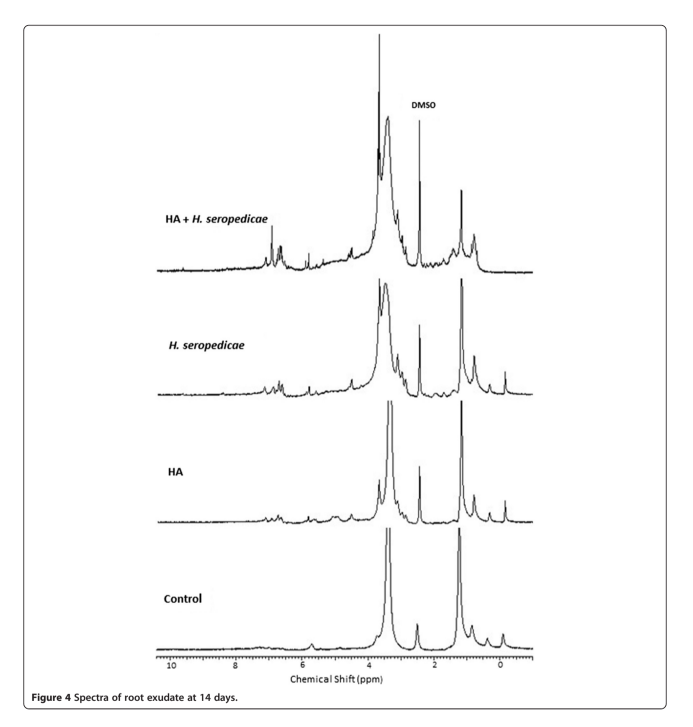
and aromatic (5.0 to 8.0 ppm) regions. NMR spectra revealed different exudate profiles between treated and control plants. On day 7, the control spectrum contained several chemical shifts in the aliphatic region, with a small signal at 0.96 ppm and a short, intense signal at 1.24 ppm. The sugar region had a main signal at 3.71 ppm and other signals of low intensity at 3.41 and 3.93 ppm. In the aromatic region, it was possible to



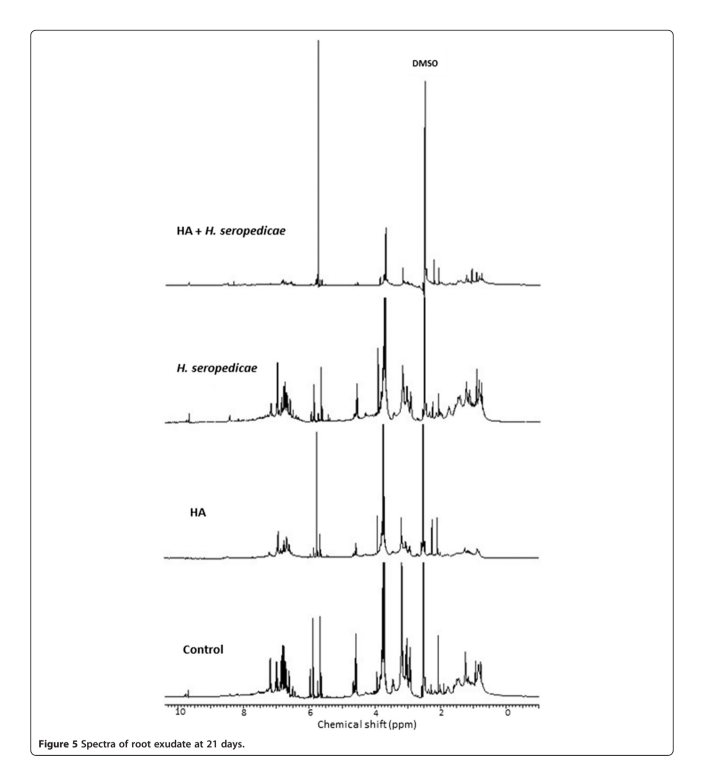


observe small signals in the 6.69 to 6.74 ppm region and at 7.0 ppm. The signals at 1.24 ppm may have been related to the presence of  $CH_3$  compounds of aliphatic acids and  $\beta CH_3$  in amino acids of maize extracts. The strong absorption at 3.71 ppm may be attributed to the presence of glycosylated compounds ( $\beta$ -Glc and  $\alpha$ -Glc). The signals at 6.7 ppm are typical of hydroxybenzoic acids, such as cinnamic and protocatechuic acids. Tryptophan, histidine, and gallic acids have signals near 7.0 ppm. Treatment of maize seedlings with HA changed the region of aliphatic absorption, and additional signals were observed at 0.84, 0.88, and 1.10 ppm. These signals were also recorded for exudates from plants treated with *H. seropedicae* (treatment PGPB), which had an additional signal at 1.64 ppm that was absent in control exudates. Additional signals were present at 1.08, 1.20, and 2.0 ppm in exudates from treatment HA + PGPB. The exudate spectrum from seedlings treated with HA alone had an additional absorption at 3.93 ppm attributable to sucrose (3.48, 3.84, 3.90, 4.22, and 5.42 ppm) and/or lysine (since a typical signal was also observed at 1.64 ppm [ $\delta$ CH<sub>2</sub>]). Exudates from seedlings in treatment HA + PGPB had additional though very small signals in the aromatic region in the range 6.34 to 6.62 ppm and a very small signal at 8.52 ppm. Compounds similar to niacin have signals in this region.

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The profiles of exudates collected on day 14 were very different from those collected on day 7. The intensity of the signal at 1.23 ppm was enhanced in all spectra, with the highest intensity in the exudates collected from seedlings in treatment PGPB. We observed very weak signals related to aromatic compounds in the control exudate spectrum on day 14, with just one visible signal at 5.70 ppm. In contrast, signals for this region were very strong in treatment HA + PGPB exudates. In treatment HA, there was a diversity of signal shoulders in regions 4.99 to 5.13, 5.65 to 5.71, and 6.69 to 6.80 ppm and individual signals at 6.99 and 7.18 ppm. Treatment PGPB exudates had clearer signals in this region than treatment HA; treatment PGPB had well-resolved signals at 5.64, 5.85, 6.07, 6.77, 6.93, and 7.20 ppm. This spectrum was marked by weak main signals at 3.54 and 3.72 ppm. There were additional signals in the aliphatic region at 1.48, 1.77, and 2.03, which were not present in the spectra from the controls and treatment HA. The characteristics of the exudate spectra were similar between



treatments PGPB and HA + PGPB, i.e., a strong modification in the aromatic region including a sharp signal at 6.98 ppm. However, in treatment HA + PGPB, there were changes in the sugar region including an increased intensity in the signal at 3.72 ppm.

On day 21 after inoculation, seedlings were less dependent on seed reserves and the exudate profiles were modified by the treatments (Figure 2, Table 1). The spectra were more complex than those in earlier exudates; they were characterized by an abundance of signals. Control exudates had a wide range of small signals in the aliphatic region. In the control spectra from days 7 and 14, the main signal in this region was at 2.08 ppm. Additional signals occurred in the sugar region between

# Table 1 1H chemical shift

7	14	21	7								
		21	7	14	21	7	14	21	7	14	21
0.96	0.10	0.10 to 0.72	0.84	0.38	0.68	0.84	0.38	0.01 to 0.59	1.08	0.77	0.78
1.24	0.38	0.77	0.93	0.85	0.70	0.93	0.85	0.62	1.10	0.85	0.82 to 0.89
	0.85	0.80 to 1.39	1.10	1.23	0.71	1.10	1.23	0.64	1.20	1.23	0.94
	1.23	1.44	1.24		0.73 to 1.00	1.24	1.48	0.65	1.24	1.49	0.95
		1.47 to 1.50	1.64		1.03 to 1.22	1.64	1.77	0.70	2.0	2.29	1.07 to 1.10
		1.54 to 1.68			1.25		2.03	0.73	3.05	2.35	1.17 to 1.28
		1.76			1.28 to 1.54			0.77	3.18	2.92 to 3.17	1.41 to 1.54
		1.78			1.57			0.80 to 0.86			2.09
		1.81			1.60 to 2.06			0.90 to 1.24			2.24
		1.84			2.09			1.34			2.46 to 2.57
		1.86 to 2.39			2.10			1.35			2.93
		2.42			2.14 to 2.22			1.39			2.95
		2.43			2.25			1.44			2.99 to 3.28
		2.47			2.28			1.48 to 1.51			
		2.52			2.30			1.54			
		2.56			2.34 to 2.42			1.57			
		2.57			2.47			1.60 to 1.69			
		2.69			2.52			1.76			
								1.77			
								1.80			
								2.52			
3.47	3.39	3.38 to 4.72		3.40	3.27 to 3.61	3.50			3.44	3.46	3.40
			3.74				3.54				3.42
											3.53
											3.56 to 3.82
											3.88
					1107 10 1155		1.05				5.00
6.69 to 6.74	5 70	5 4 5	671	5.65 to 5.71	5.00 to 6.16	651	5 64	5.01 to 6.80	634 to 662		5.66
	5.7 0										5.72
											5.77
											5.82
											6.82
				7.10							6.83
											8.34
							7.20				0.54
								9.09 10 9.99		7.10	
					9.74						
	3.47 3.71 3.93 6.69 to 6.74 7.0	<ul> <li>1.23</li> <li>3.47</li> <li>3.71</li> <li>3.73</li> <li>3.93</li> <li>6.69 to 6.74</li> <li>5.70</li> </ul>	<ul> <li>1.23</li> <li>1.44</li> <li>1.47 to 1.50</li> <li>1.54 to 1.68</li> <li>1.76</li> <li>1.78</li> <li>1.81</li> <li>1.84</li> <li>1.86 to 2.39</li> <li>2.42</li> <li>2.43</li> <li>2.47</li> <li>2.52</li> <li>2.56</li> <li>2.57</li> <li>2.69</li> <li>2.75 to 3.36</li> </ul> 3.47 <ul> <li>3.47</li> <li>3.39</li> <li>3.38 to 4.72</li> <li>3.73</li> <li>4.92</li> <li>3.93</li> </ul> 6.69 to 6.7 <ul> <li>5.70</li> <li>5.45</li> </ul>	1.23 1.44 1.47 to 1.00 1.47 to 1.00 1.47 to 1.00 1.76 1.78 1.78 1.81 1.84 1.84 1.84 1.84 2.42 2.43 2.47 2.52 2.56 2.57 2.56 2.57 2.69 2.75 to 3.36 3.71 3.73 3.71 3.79 3.74 3.71 3.79 3.74 3.79 3.74 3.71 3.74 3.74 3.74 3.74 3.75 5.70 5.71 5.70 5.71 5.70 5.71 5.71 5.71 5.71 5.71 5.71 5.71 5.71	1.23 1.44 1.47 to 1.50 1.54 1.47 to 1.64 1.54 to 1.68 1.76 1.78 1.81 1.84 1.84 1.84 1.84 1.84 1.85 to 2.39 2.42 2.43 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      2.34 to 2.42         2.52       2.47       2.52       2.30       2.56       2.57       2.67       2.66       2.57       2.67       2.66       2.57       2.67       2.67       2.57       2.67       2.57       2.67       2.57       2.67       2.57       2.67       2.57       2.67       2.57       2.67       2.57       2.67       2.57       2.67       2.57       2.67       2.57       2.67       2.57       2.67       2.57       2.67       2.57       2.67       2.57       2.57       2.67       2.57       2.57       2.67       2.57       2.66       3.67       3.68       3.74       3.68       3.67       3.68       3.67       3.66       3.69       3.67       3.66       3.66       3.66       4.57       3.66       4.56       4.56       4.56 <td>1231.441.240.73 to 1.001.241.481.47 to 1.001.641.03 to 1.221.641.711.54 to 1.681.25 to 1.681.25 to 1.681.571.761.28 to 1.511.571.571.571.811.60 to 2.061.572.160 to 2.061.571.842.102.10 to 2.262.14 to 2.221.572.622.242.262.301.572.522.34 to 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3.03 and 3.45 ppm, including a sharp signal at 3.18 ppm. The main absorption in this region occurred at 3.73 ppm, as in earlier exudates. Signals between 4.55 and 4.67 ppm were very minor or absent in exudates from days 7 and 14, but much higher and with greater definition by day 21, which was also the case for signals in the aromatic region (6.50 to 7.19 ppm). Additional small signals occurred at 9.69, 9.76, and 9.78 ppm. The exudate profile from treatment HA was generally quite similar to that of the control, but significant differences were apparent. For example, there was no sharp, high-intensity signal at 2.25 ppm in the control, but this was observable in treatment HA exudates, which also had a very intense signal at 5.57 ppm (this was the main signal in treatment HA + PGPB spectra but a very minor one in treatment PGPB and in the control). On day 21, the spectra of exudates were closely similar between treatment PGPB and the control.

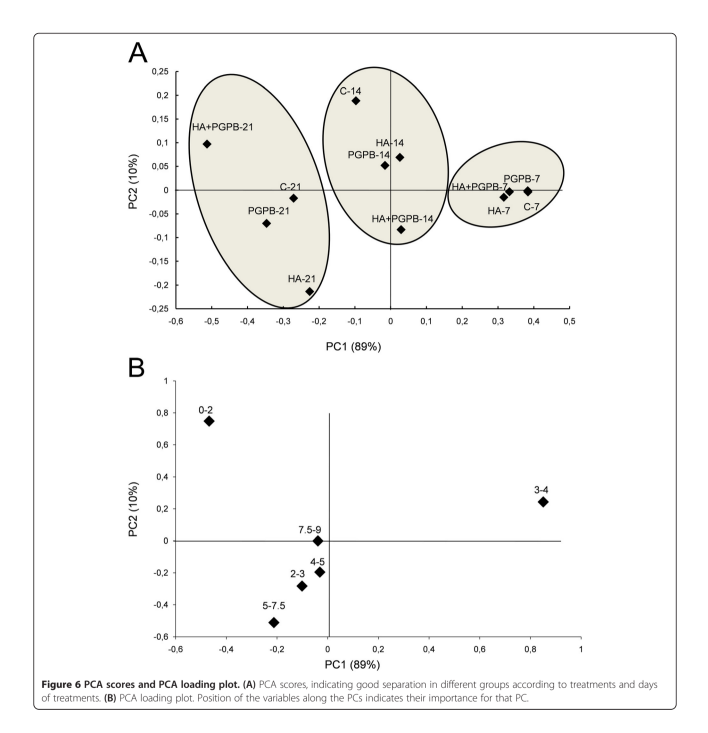
Similarities and dissimilarities among treatments and experimental days were examined by PCA analysis (Figure 6). PC1 (89%) and PC2 (10%) accounted for 99% of the total variance in the <sup>1</sup>H NMR data set. Symbols in the PCA plot are identified by treatment codes and days of exudate collection. Three groups of symbols are apparent in the Cartesian two-dimensional space of axes PC1 and PC2; these correspond to the three occasions of exudate collection. The day 7 group is located in the positive sector of the PC1 axis and in the negative sector of the PC2 axis. Positive values of PC1 and the loading coefficient responsible for this distinction are large for the carbohydrate <sup>1</sup>H NMR region (3 to 4 ppm). HA + PGPB symbols for the day 14 fall in the negative sector of PC2, but other treatment symbols for this collection day fall in the positive sector of this axis. HA and PGPB symbols cluster together; the control treatment had high values of aliphatic moieties (0 to 2 ppm). On day 21, the HA + PGPB and HA treatments had very different exudate compositions with elevated aromatic and aliphatic contents, respectively. The control and PGPB treatment clustered together.

Overall, the PCA analysis aided interpretation and comparisons in the data set. Thus, on day 7, exudate compositions were not significantly different among treatments, but significant differences were observed on days 14 and 21. This trend was corroborated by GC-MS analysis. The chromatograms are available in the supplementary data (Additional files 1, 2, and 3). Main exudates retained in the RP C18 column and identified by GC-MS are indicated in Table 2. On day 7, the chromatograms were closely similar; the main compounds were nitrogenous carbohydrate derivatives and fatty acids (as methyl esters). Nevertheless, there were some differences among treatments on day 7. For example, exudation of carbohydrate compounds by root seedlings in treatment HA + PGPB exceeds those in other treatments (Table 2). The chromatogram peaks of these carbohydrate compounds for treatments PGPB and HA were twice as high as that for the control; we identified galacto, gluco, and mannose pyranoside structures in these peaks. D-glucopyranoside moieties associated with aromatic rings were also found in control exudates, but 1,6-β-glucose was found only in exudates from plants in treatment HA. The quantities and diversities of nitrogenous compounds on day 7 were also greater in treatment HA and PGPB exudates than in the controls (Table 2). Benzilamines, polyamines, pyrrol derivatives, amino acid complexes, and nucleotide derivatives were included in this group of compounds. Benzilamines and polyamines were exuded only in treatments HA and HA + PGPB. Nitrogenous compounds from the phenylalanine pathway were present; these included pyrimidinedione, pyrrole-3-carboxylic acid and a derivative from *p*-coumaric hydrolysis with a retention time at 28.91 min (identified as caffeine). Structures derived from adenosine were found in all treatments, but those from purine were found only in treatment PGPB. Overall, treatment HA + PGPB induced the largest and most diverse exudation of nitrogenous compounds.

Fatty acids comprised the second largest class of compounds exuded on day 7. Unlike nitrogenous compounds, treatments PGPB and HA + PGPB induced exudations of only small quantities of fatty acids (in comparison with treatment HA and the control). Chromatogram peak areas for fatty acids were elevated by 52% in treatment HA in comparison with the control on day 7. The main fatty acid in the HA exudate was hexadecanoic acid (palmitic acid). Stearic acid (octadecenoic) was exuded only by control plants, while tridecanoic acid and eicosanoic acid (also known as arachidic acid), which are minor constituents of corn oil, were found only in treatment HA exudates. Palmitic and stearic acids were found in treatment PGPB exudates. Although GC-MS is not particularly appropriate for short-chain organic acid identification, we did observe the presence of propenoic acid associated with long-chain carbon compounds in control and treatment HA exudates; we also detected succinic acid associated with aromatic moieties in treatment HA + PGPB. Benzenedicarboxylic acid was the main aromatic exudate. The steroid isosteviol was found in all exudates. Other steroids, such as cyclolanostane and androstane, were found in exudates produced in treatments HA, PGPB, and HA + PGPB.

The main classes of compounds were similar among days 7, 14, and 21 (Table 2), but the diversity of compounds diminished progressively as the seedlings grew. There were differences among the treatments: on day 14, nitrogenous compounds were still abundant in the HA, PGPB, and HA + PGPB treatment exudates; the concentrations were threefold to fourfold higher than those in the control. Nucleotide derivatives were observed on day 7, but not later. Products from phenylalanine and pyridines were exuded in treatments PGPB and HA + PGPB,

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while phenyl cyanate and pyrrol derivatives were found in treatment HA exudates. The compounds present in exudates from treatments PGPB and HA + PGPB were closely similar (Table 2), although benzene, 1-isocyanato-4-methoxy, was exuded only in treatment PGPB and carbonic acid, monoamine, *N*-(2,4-dimethoxyphenyl)-, and butyl ester only in treatment HA + PGPB. Exudations of carbohydrates were elevated in treatments HA and PGPB. These exudates contained compounds we had identified on day 7. However, quantities of carbohydrate derivatives were reduced in treatment HA + PGPB and absent in control seedling exudates.

Hexadecanoic and octadecanoic acids were the main fatty acids occurring as methyl esters in control exudates. These compounds also occurred in other treatments, but the diversity was greatest in PGPB and HA + PGPB treatments. However, on day 7, the quantities and diversity were highest in treatment HA. As on day 7, products of the isosteviol reaction occurred in all exudates; another steroid (androstane derivative) was also

# Table 2 Compounds identified by GC-MS analysis

Retention time (min)	Compound class		Control			HA			H. seropedicae			HA + H. seropedicae		
		7	14	21	7	14	21	7	14	21	7	14	21	
	Nitrogenous compounds													
9.575	3-Butenoic acid, 3-methoxy-4-nitro-, methyl ester, (E)	nd	nd	nd	nd	3.82	nd	nd	nd	nd	nd	nd	nd	
14.754	Benzene, 1-isocyanato-4-methoxy	nd	nd	nd	nd	0.82	nd	nd	0.83	nd	0	nd	nd	
20.618	3,4-Dimethoxy-6-amino toluene	8.54	2.75	0	8.93	4.78	6.08	7.41	5.00	9.70	12.14	5.43	3.35	
21.787	1 <i>H</i> -Pyrrole-3-carboxylic acid, 5-formyl-1-(methoxymethyl)-, methyl ester	4.22	nd	nd	nd	1.21	nd	nd	nd	2.65	nd	nd	nd	
21.795	3,4,5-Trimethoxybenzylamine	nd	nd	nd	5.11	nd	nd	3.47	1.18	1.18	4.56	1.21	nd	
24.873	2,5-Dimethoxyphenyl isocyanate	11.41	7.41	0	16.78	13.18	24.30	14.36	10.64	48.74	18.82	24.19	16.9	
25.952	2,4-Dimethoxyphenylamine, N-ethoxycarbonyl-	0.48	nd	nd	0	nd	nd	nd	0.26	0.57	0.68	0.70	nd	
26.019	9H-Purin-6-amine, N,N,9-trimethyl	2.15	nd	nd	1.30	nd	nd	1.97	nd	nd	1.84	nd	nd	
26.982	4-(Diethylamino)salicylaldehyde	nd	nd	nd	nd	nd	nd	nd	nd	2.87	nd	nd	nd	
27.134	2,3,4-Trimethoxyphenylacetonitrile	nd	nd	nd	nd	nd	nd	nd	nd	0.94	nd	nd	nd	
28.503	Bicyclo[2.2.1]heptane-2-acetamide, N-(1,3-dihydro-5,6-dimethoxy-3-oxo- 4-isobenzofuranyl)	3.04	nd	nd	4.69	nd	5.30	3.51	nd	13.17	5.26	nd	3.32	
28.703	3,4,5-Trimethoxyphenyl cyanate	nd	nd	nd	nd	2.23	nd	nd	nd	nd	nd	nd	nd	
28.709	Tris(2,4,6-dimethylamino)pyrimidine	nd	nd	nd	nd	nd	nd	nd	1.75	nd	nd	3.34	nd	
28.915	Caffeine	nd	nd	nd	0.48	nd	nd	nd	nd	nd	nd	nd	nd	
29.340	Acetamide, N-[2-(3,4-dimethoxyphenyl)ethyl]	nd	nd	nd	nd	nd	nd	nd	0.89	4.06	nd	nd	nd	
29.346	3-Pyridinecarboxamide, 4-(methoxymethyl)- 6-methyl-2-propoxy-	nd	nd	nd	nd	nd	0.53	nd	nd	nd	nd	nd	1.49	
29.537	1,3-Benzenediamine, N,N'-diethyl	nd	nd	nd	nd	0.41	nd	nd	nd	nd	nd	nd	nd	
30.534	Carbonic acid, monoamide, N-(2,4-dimethoxyphenyl)-, butyl ester	nd	nd	9.55	nd	nd	nd	nd	nd	nd	nd	nd	nd	
30.774	3-(2-Ethoxycarbonyl-ethyl)-4-methyl-1 <i>H</i> -pyrrole- 2-carboxylic acid, ethyl ester	nd	nd	nd	nd	12.62	nd	nd	nd	nd	nd	nd	nd	
30.775	Carbonic acid, monoamide, N-(2,4-dimethoxyphenyl)-, butyl ester	nd	nd	nd	nd	nd	nd	nd	nd	nd	nd	6.93	nd	
30.801	Propanamide, 3-amino-N-(2,5-dimethoxyphenyl)- 3-(hydroxyimino)-	nd	nd	nd	nd	nd	nd	nd	14.99	nd	nd	nd	nd	
26.134	Phenol, 4-(aminomethyl)-2-methoxy	nd	nd	nd	nd	0.33	nd	nd	nd	nd	nd	nd	nd	
30.741	1,5-Bis[veratrylaminomethyl]-2,6- dihydroxynaphthalene	nd	1.25	nd	nd	nd	nd	nd	nd	nd	nd	nd	nd	
30.952	3H-Purin-6-amine, N,N,3-trimethyl	nd	nd	nd	nd	nd	nd	0.61	nd	nd	0.58	nd	nd	
31.834	Benzeneethanamine, N-acetyl-2,3,4,5-tetramethoxy	nd	nd	nd	nd	nd	nd	nd	2.83	nd	nd	nd	nd	
32.132	Acetamide, N-(3,4,5-trimethoxyphenethyl)	nd	nd	nd	nd	nd	nd	nd	nd	0.91	nd	nd	0.40	
35.077	Uridine, 2'-deoxy-3-methyl-3',5'-di-O-methyl-	nd	nd	nd	nd	nd	nd	nd	nd	nd	0.19	nd	nd	
35.926	Cytidine, N-methyl-, 2',3',5'-trimethyl ether	0.62	nd	nd	1.44	nd	nd	1.52	nd	nd	0.55	nd	nd	
36.019	Thymidine, 2'-deoxy-N,O,O-trimethyl-	0.91	nd	nd	1.62	nd	nd	2.24	nd	nd	3.37	nd	nd	
37.792	N-[12-Aminododecyl]aziridine	0.52	nd	nd	0.58	nd	nd	0	nd	nd	0.95	nd	nd	
41.125	Adenosine, 2'-deoxy-N,N,O,O-tetramethyl-	nd	nd	nd	0.65	nd	nd	1.60	nd	nd	2.11	nd	nd	
42.007	Adenosine, 3'-amino-3'-deoxy-N,N-dimethyl	nd	nd	nd	0.20	nd	nd	0.46	nd	nd	0.66	nd	nd	
		nd	nd	nd	nd	nd	nd	nd	nd	nd	nd	nd	nd	

# Table 2 Compounds identified by GC-MS analysis (Continued)

	Carbohydrate compounds												
17.168	1,6-Anhydro-β-D-glucose, trimethyl ether	nd	nd	nd	1.35	nd	nd	nd	0.77	nd	0.83	nd	nd
18	α-D-Galactopyranoside, methyl 2,3,4,6-tetra-O-methyl	1.74	nd	nd	3.17	nd	nd	0.73	nd	nd	4.24	nd	nd
17.688	3,4,6-Tri-O-methyl-D-glucose	nd	nd	nd	nd	0.54	nd	nd	0.83	nd	nd	nd	nd
18.1280	a-Methyl 4-methylmannoside	nd	nd	nd	nd	nd	nd	nd	0.55	nd	nd	nd	nd
21.355	2,4,5,6,7-Pentamethoxyheptanoic acid, methyl ester	nd	nd	nd	nd	0.33	nd	nd	nd	nd	nd	nd	nd
20.206	β-D-Glucopyranose, 1,6-anhydro	nd	nd	nd	nd	nd	nd	nd	nd	1.02	nd	nd	nd
28.814	1,2,3,4-Tetramethylmannose	nd	nd	nd	nd	1.85	nd	nd	1.75	nd	nd	0.58	nd
28.631	α-D-Mannopyranoside, methyl 2,3,4,6-tetra-O-methyl	nd	nd	3.33	nd	nd	nd	0	nd	nd	nd	nd	0.53
37.425	α-D-Glucopyranoside, phenyl 2, 3,4,6-tetra-O-methyl	nd	nd	nd	nd	41.61	31.93	nd	22.57	nd	nd	5.62	nd
	Fatty acids as methyl ester												
26.320	Tridecanoic acid, 12-methyl-, methyl ester	0.77	nd	nd	0	nd	nd	1.06	1.26	nd	0.80	1.62	nd
26.323	Methyl tetradecanoate	nd	nd	nd	1.22	nd	nd	nd	nd	nd	0	nd	nd
27.419	2-Hexadecenoic acid, methyl ester, (E)	nd	nd	nd	0.32	nd	nd	nd	nd	nd	0.37	nd	nd
27.901	Tridecanoic acid, 12-methyl-, methyl ester	nd	nd	nd	0.28	nd	nd	nd	nd	nd	0	nd	nd
30.133	Methyl hexadec-9-enoate	1.06	nd	nd	2.94	nd	nd	0.84	nd	nd	0.47	nd	nd
30.181	9-Octadecenoic acid (Z)-, methyl ester	0	nd	nd	0	nd	nd	nd	nd	nd	0.13	nd	nd
30.625	Hexadecanoic acid, methyl ester	4.22	2.26	0	7.28	nd	nd	4.05	nd	1.04	3.22	3.31	1.96
30.327	6-Octadecenoic acid, methyl ester	nd	nd	nd	nd	nd	nd	nd	nd	nd	nd	0.82	nd
34.053	9-Octadecenoic acid, methyl ester, (E)	1.27	nd	nd	nd	nd	nd	nd	0.73	nd	nd	0.84	nd
34.164	8-Octadecenoic acid, methyl ester, (E)	0.67	nd	nd	nd	nd	nd	nd	nd	nd	nd	nd	nd
34.548	Octadecanoic acid, methyl ester	0.95	1.11	0	1.36	0.57	nd	2.15	1.61	0.28	1.40	1.60	0.67
38.161	Eicosanoic acid, methyl ester	nd	nd	nd	0.16	nd	nd	nd	nd	nd	nd	nd	nd
	Organic acids												
26.507	Benzoic acid, 4,5-dimethoxy-2- (4-methoxybenzenesulfonylamino)-	nd	nd	nd	nd	1.31	nd	nd	nd	nd	nd	nd	nd
40.052	Succinic acid, di(5-methoxy-3- methylpent-2-yl) ester	nd	nd	nd	nd	nd	nd	nd	nd	nd	0.36	nd	nd
43.743	Cyclopropanecarboxylic acid, 2,2-dimethyl-3-(2-methyl-1-propenyl)-, 2-methyl-4-oxo-3-(2-propenyl)-2- cyclopenten-1-yl ester	nd	nd	nd	nd	nd	nd	nd	0.70	nd	nd	nd	nd
	Aromatics and phenol derivatives												
15.817	2-Methoxy-4-vinylphenol	nd	nd	nd	nd	nd	nd	nd	nd	0.3	0	nd	nd
19.337	Phenol, 4,6-di(1,1-dimethylethyl)-2-methyl	nd	nd	nd	0.24	nd	nd	nd	nd	nd	0.23	nd	nd
24.653	Isoelemicin	nd	nd	nd	nd	nd	nd	nd	nd	0.49	0	nd	nd
37.592	Benzene, (1-methyldodecyl)	0.31	nd	nd	nd	nd	nd	1.98	nd	nd	0.80	nd	nd
41.779	1,2-Benzenedicarboxylic acid, diisooctyl ester	nd	nd	69.71	nd	nd	nd	1.49	nd	nd	0.44	nd	nd
41.784	1,2-Benzenedicarboxylic acid, mono(2-ethylhexyl) ester	1.55	nd	nd	nd	nd	2.37	nd	nd	nd	0	nd	7.49
21.512	Phenol, 2,6-dimethoxy-4-(2-propenyl)	nd	nd	nd	nd	0.25	nd	nd	nd	nd	nd	nd	nd
35.007	Phenol, 5-methoxy-2,3,4-trimethyl	nd	0	nd	nd	0.22	nd	nd	nd	nd	nd	nd	nd
37.803	Benzene, (1-methylnonadecyl)	nd	0.78	nd	nd	nd	nd	nd	1.15	nd	nd	1.36	nd

# Table 2 Compounds identified by GC-MS analysis (Continued)

22.015	Benzoic acid, 4-hydroxy-3,5-dimethoxy	nd	nd	nd	nd	0.21	nd	nd	nd	nd	nd	nd	nd
22.016	1,2,3,4-Tetramethoxybenzene	nd	nd	0.27	nd								
23.399	Benzoic acid, 3,4-dimethoxy-, methyl ester	nd	nd	nd	nd	0.72	1.41	nd	nd	nd	nd	nd	1.65
25.152	2-Propenoic acid, 3-(4-methoxyphenyl)-, methyl ester	nd	nd	nd	1.17	1.03	nd	nd	0.65	nd	nd	0.66	nd
26.277	Benzenepropanoic acid, 3,4-dimethoxy-, methyl ester	nd	nd	nd	nd	nd	5.14	nd	nd	nd	nd	nd	3.27
26.328	Benzoic acid, 3,4,5-trimethoxy-, methyl ester	nd	nd	nd	0	nd	nd	nd	nd	2.70	nd	nd	nd
29.738	2-Propenoic acid, 3-(3,4-dimethoxyphenyl)-, methyl ester	0.49	nd	nd	0	2.61	5.02	nd	0.75	4.24	0.23	1.24	3.21
36.154	2-Propenoic acid, 2-methyl-, dodecyl ester	0	nd	nd	0	nd	nd	nd	nd	nd	0.12	nd	nd
	Terpenoids and steroids												
43.074	17β-Hydroxy-6α-pentyl-4-oxa-5β-androstan-3-one	nd	nd	nd	0.74	nd	nd	0.93	nd	nd	nd	nd	nd
43.412	Dihydroxyisosteviol	3.88	nd	nd	3.94	nd	nd	5.45	3.95	nd	3.69	0.73	nd
42.582	Dihydro-isosteviol methyl ester	nd	2.04	nd	nd	0	9.42	nd	nd	nd	nd	nd	17.03
43.332	Methyl dihydroisosteviol	nd	nd	nd	nd	1.47	nd	nd	nd	nd	nd	nd	nd
43.665	Isosteviol methyl ester	nd	2.88	nd	nd	4.81	nd	nd	4.90	17.18	nd	4.68	nd
39.849	5-Cholene, 3,24-dihydroxy-	nd	0.12	nd	nd								
43.430	Patchoulene	nd	nd	nd	0.36	nd	nd	nd	nd	nd	nd	nd	nd
45.538	Androstan-17-one, 3-ethyl-3-hydroxy-, (5α)	nd	0.78	nd	nd	2.89	nd	nd	1.03	0.50	nd	2.30	nd
42.354	5α,6α-Epoxy-17-oxo-6β-pentyl-4-nor-3, 5-secoandrostan-3-oicacid, methyl ester	nd	nd	nd	0.31								
43.091	5β-Pregnan-17α,21-diol-3,20-dione	nd	nd	nd	25.88								
43.434	Pregnan-20-one, (5a,17a)	nd	nd	2.48	nd	nd	nd	nd	nd	nd	nd	nd	nd
	Alcohols												
3.974	Ethanol, 2-(2-methoxyethoxy)	nd	1.18	0.33	nd	nd							
6.715	1-Hexanol, 2-ethyl	1.65		7.48	2.71	0.39	1.13	0.86	nd	nd	3.39	nd	1.19
12.925	Ethanol, 2-phenoxy	nd	0.63	nd	nd	nd							
42.010	9-t-Butyltricyclo[4.2.1.1(2,5)]decane-9,10-diol	nd	nd	1.03	nd								
	Unidentified												
30.542		nd	nd	nd	3.14								
31.625		nd	nd	nd	0.74								
31.825		nd	nd	nd	nd								
32.184		nd	nd	nd	nd	0.79	nd	nd	nd	nd	nd	nd	nd
35.248		nd	nd	nd	nd	0.77	nd	nd	nd	nd	nd	nd	nd
38.362		nd	0.42	nd	nd								
39.253		nd	0.09	nd	nd								
40.033		nd	nd	0.43	nd								
40.230		nd	nd	nd	0.20	nd	nd	nd	nd	nd	0.37	nd	nd
40.833		nd	nd	nd	0.18								
40.964		nd	0.31	nd	nd								
41.908		nd	nd	nd	0.49	nd	nd	nd	nd	nd	0.41	nd	nd
42.157		nd	nd	0.24	nd								
43.746		nd	0.43	nd	0	nd	nd						
44.305			nd		nd							nd	

44.532	nd	nd	nd	0	nd	nd	nd	nd	nd	nd	nd	0.45
45.275	nd	nd	nd	0.74	nd	nd	0.35	nd	nd	0.89	nd	nd
46.026	nd	nd	nd	nd	nd	nd	0	nd	nd	0.11	nd	nd
46.277	nd	nd	nd	nd	nd	nd	0	nd	nd	nd	nd	1.72
46.606	nd	nd	nd	nd	nd	0.53	0	nd	nd	0	nd	nd

Table 2 Compounds identified by GC-MS analysis (Continued)

Compound exudates from maize seedlings at different growth stages (7, 14, and 21 days) induced or not (control) by humic acids (HA), Herbaspirillum seropedicae (H. seropedicae), and HA and H. seropedicae. nd, not detected.

found in the exudates, though in only small quantities in the control.

Compounds identified on day 21 are shown in Table 2. Exudates collected on that day contained a smaller diversity of compounds than earlier collections. Hexanol was found in control exudates and ethanol phenoxy products in treatment PGPB, likely due to carbohydrate hydrolysis reactions. Nitrogenous compounds were the main exudate products retained by the RP C18 column; carbonic acid, monoamide, N-(2,4-dimethoxyphenyl)-, and butyl ester were found only in control exudates on day 21. 2,5-Dimethoxyphenyl isocyanate occurred in elevated quantities in exudates of treatments HA, PGPB, and HA + PGPB; a derivative from acetamide and dimethoxy amino toluene was also abundant. A nitrogenous compound identified as 4-(diethylamino)salicylaldehyde was found in treatment PGPB exudates on day 21, but not on days 7 and 14. Finally, products from acetamide were typical of exudates in treatments PGPB and HA + PGPB; they were found on day 21 at 29.34 and 32.13 min.

## Discussion

Our aim in the present work was to determine the effects of single and combined applications of humic acid and the endophytic diazotrophic bacterium *H. seropedicae* on exudation profiles of maize roots. This was a first step in identifying compounds that may be involved in rhizosphere interactions and modulation of root colonization and endophytic establishment of bacteria.

We demonstrated enhanced root colonization by *H. seropedicae* in the presence of humic acids (Figure 1). Previous work showed that the elevated bacterial population associated with maize roots may be explainable in part by humic acid sorption at the plant cell wall surface, which is associated with increased attachment and endophytic colonization by *H. seropedicae* [9,10]. Moreover, the main infection sites for *H. seropedicae* in grasses are the points of lateral root emergence, which are induced by humic acid and may contribute to enhanced establishment of the endophytic population [11].

Canellas et al. [12] and Puglisi et al. [13] previously observed changes in exudation profiles of organic acids in maize seedlings treated with humic acid. However, identifying causal relationships in changed exudate profiles is complex, and results may not always be directly attributable to treatment effects on the quantities and the composition of root exudates. Juo and Storzky [36] observed decreases in protein and carbohydrate release with increasing plant age using an electrophoretic approach. They suggested that the RP C18 column used to retain the exudates may have limited their quantitative analysis. We noticed on day 21 (across all treatments) that exudate yields had declined (Figure 2) relative to earlier collections on days 7 and 14.

<sup>1</sup>H NMR spectroscopy (Figures 3,4,5) demonstrated differences in exudate profiles under the influence of the treatments, and PCA analysis revealed significant changes from day 14 onward (Figure 6). Roncatto-Maccari et al. [37] observed that 3 days after maize inoculation, the population of *H. seropedicae* mainly comprises single cells attached to the root surface. They (*loc. cit.*) found that after 12 days the cells were connected and forming aggregates; they detected (by scanning electron microscopy) a halo of mucilage surrounding the bacterial colony on the roots of inoculated maize. Although plant-bacterial signaling is considered a rapid phenomenon, changes in exudate NMR spectra in our experiment occurred only after extensive colonization on the root axis (i.e., on day 14).

One group of candidate biomolecules emerging from this prospective study comprised a number of compounds that have been identified as possible inducing agents of the quorum sensing (QS) system, which is a mechanism responsible for modulating population density, biofilm formation, and the specific genetic machinery for niche persistence and colonization at the root level. These compounds include oligopeptides and substituted gamma-butyrolactones, 3-hydroxypalmitic acid methyl ester, 3,4-dihydroxy-2-heptylquinoline, and a furanosyl borate diester ([5] and references therein). Interestingly, our analyses of the overall transcriptome profile of H. seropedicae strain Smr1 (as part of a broad examination of gene expression in planktonic and biofilm lifestyles) have shown that biofilm maturation is coupled with differentially regulated genes involved in aromatic metabolism and multidrug transport efflux activation (data not shown). These potential quorum sensing inducers may be key molecules in the well-known rhizosphere competence of H. seropedicae [14,25]. Functional studies linking these compounds to chemotaxis and biofilm induction are urgently required.

Another interesting future study that may develop from our prospective explorations might be an evaluation of the ability of *H. seropedicae* to use secondary metabolites as carbon sources that are exclusively or primarily exuded from maize roots treated with this bacterium. The presence of a catabolic machinery for specific cleavage of these compounds (especially compounds with antimicrobial properties) would represent a nutritional advantage for the colonization of the host plant rhizosphere.

We identified hexadecanoic acid (palmitic acid) in exudates of all treatments on days 7 and 14, but by day 21, these compounds were detected only in treatment PGPB. Non-quinoline derivatives of other heterocyclic aromatic N compounds belonging to the pyrimidine class were collected exclusively in treatments PGPB and HA + PGPB (Table 2). The most remarkable difference in root exudates among treatments was in N compounds; these increased with time in all treatments other than the controls, in which quantities decreased. Some nitrogenous compounds, like benzenamine ( $C_6H_5CONH_2$ ), 2,4-dimethoxy, 3,4-dimethoxy-6-amino toluene, and 2,5dimethoxyphenyl isocyanate, were found in the exudates; they are probably derivatives of aromatic/phenolic compounds that react with TMAH used in methoxylation since the phenolic compounds, which are the main secondary metabolites synthesized by maize, react promptly with NH<sub>2</sub>. Pyrimidinediones comprise a class of chemical compounds characterized by a pyrimidine ring substituted with two carbonyl groups. 2,4(1H,3H)-pyrimidinedione, 1,3,5-trimethyl is typical of uracil derivatives, i.e., nucleobases in the nucleic acid of RNA. Pyrrole carboxylic acid derivatives, which may originate from the L-tryptophan pathway, have been described as antibacterial agents. Finally, we found one polyamine derivative of spermine. Such derivatives have been implicated in a wide range of metabolic processes in plants, ranging from cell division and organogenesis to protection against stress; they act as plant and microbial growth stimulants and chemoattractants [5].

Stearic and palmitic acids were the main fatty acids exuded by maize seedling roots. These fatty acids are predominant in most plants; they are synthesized by acetyl-CoA carboxylase and fatty acid synthase. Quantities of these fatty acids exuded by the controls differed between days 14 and 21. Stearic and palmitic acids were exuded by maize seedlings treated with *H. seropedicae* with or without HA (Tables 2). Terpenoids derived from the kaurenoic acid pathway, including gibberellic acids, were found in all treatments. However, their diversity was highest in treatments PGPB and HA + PGPB. A significant increase in transcript levels of *ent-kaurene*  oxidase genes that are involved in gibberellin synthesis occurs in maize seedlings treated with *H. seropedicae* [38]. The agreement between gene expression in the gibberellin biosynthesis pathway (Zmko1) and the exudation profile in the same biological model supports our suggestion that the gibberellin synthesis pathway in root tissues is modulated by *H. seropedicae* colonization.

Root colonization by *H. seropedicae* in graminaceous plants requires movement of the bacteria to the root surface followed in sequence by adsorption and attachment to the cell wall surface [25,27]. Attached bacteria may persist at the root surface as aggregates or biofilms, and they occasionally trigger plant responses that induce lateral root sites, which serve as entryways for endophytic establishment in the plant host. Humic acids are heterogeneous, irregular, and amphiphilic structures with high charge densities and hydrophobic cores that facilitate adsorption phenomena and lateral root induction [39]. As *H. seropedicae* is a flagellated cell, there is a potential for induction of flagellar activity by plant-released compounds (chemotactic substances), as previously described for other bacteria [40].

From the inventory of exuded compounds, it may be possible to select candidate molecules as quorum sensing mediators of biofilm maturation, as substrates for catabolic mechanisms to supply energy, or as detoxifiers, thereby increasing the rhizosphere competence of *Herbaspirillum*. Furthermore, changes in rhizosphere hydrophobicity by fatty acid exudates may be an important mechanism for preserving cell viability. We detected relatively large amounts of gibberellic acid precursor in exudates isolated from maize seedlings treated with *H. seropedicae*; this bacterium induced changes in nitrogenous compounds as well.

#### Conclusion

The number of unidentified compounds was much higher in treatments PGPB and HA + PGPB than in the control. The limitations of a conventional GC-MS approach, such as low mass resolution and the need for previous volatilization of compounds, suggest that the instrument should be combined with other analytical tools, like liquid chromatography coupled to high mass definition spectroscopy. We found that root exudates from maize seedlings mostly comprised fatty acids and nitrogenated compounds. We observed changes in root exudate profiles in maize seedlings treated with HA and *H. seropedicae*. These changes were most pronounced on days 14 and 21, when there was enhanced fatty acid exudation in treatment HA and elevated nitrogenate and terpene exudation from treatment PGPB.

This is the first study to date that has evaluated the combined application of a plant growth-promoting bacteria and humic acid on maize root exudation profiles. This study was carried out on controlled conditions in a hydroponic culture. Overall, the chemical changes we observed in the rhizosphere may improve the survival, persistence, and biological activity of *H. seropedicae* in the presence of humic substances and provide a pool of candidate molecules for use as additives in inoculant technology formulation, which will contribute substantially to progress in arable agriculture. However, the findings of this hydroponic study need to be confirmed under real soil conditions.

# **Additional files**

Additional file 1: Figure S1. GC of root exudate at 7 days. Additional file 2: Figure S2. GC of root exudate at 14 days. Additional file 3: Figure S3. GC of root exudate at 21 days.

#### Competing interests

The authors declare that they have no competing interests.

#### Authors' contributions

LSL conducted the laboratory experiments and did the first inventory of exudates. FLO and LPC wrote and discussed the manuscript. RRO performed and interpreted the NMR data. MRGV reviewed the nomenclature of compounds and the biosynthesis route. NOA was responsible for the multivariate analysis. All authors read and approved the final manuscript.

#### Acknowledgements

The work was supported by Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq), Fundação de Amparo à Pesquisa do Estado do Rio de Janeiro (FAPERJ), Instituto Nacional de Ciência e Tecnologia (INCT) para a Fixação Biológica de Nitrogênio, Internacional Foundation of Science (IFS), and OCWP.

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#### Received: 12 June 2014 Accepted: 30 October 2014 Published online: 27 November 2014

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#### doi:10.1186/s40538-014-0023-z

**Cite this article as:** da Silva Lima *et al.*: **Root exudate profiling of maize seedlings inoculated with** *Herbaspirillum seropedicae* and humic acids. *Chemical and Biological Technologies in Agriculture* 2014 1:23.

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