1	Title: Association between gentamicin resistance and stress tolerance in water isolates of
2	Ralstonia pickettii and R. mannitolilytica
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29 ABSTRACT

Members of the species Ralstonia pickettii and R. mannitolilytica, although ubiquitous and 30 lacking major virulence factors, have been associated with nosocomial outbreaks. 31 32 Tolerance to metals, antibiotics and disinfectants may represent an advantage for their 33 ubiquity and opportunistic pathogenic potential. In this study we compared five strains that differed on the origin (hospital effluent, tap water, mineral water) and in the susceptibility 34 to aminoglycosides, regarding their tolerance to metals and disinfection. The growth 35 36 kinetics and biofilm formation capacity were tested in four R. pickettii strains and one R. *mannitolilytica* at sub-inhibitory concentrations of aminoglycosides or arsenite. The 37 survival to UV radiation, chlorine or hydrogen peroxide was also compared in 38 aminoglycoside resistant and susceptible strains. Aminoglycoside resistant strains presented 39 a higher tolerance to arsenite than the susceptible ones and either aminoglycosides or 40 arsenite were observed to stimulate the biofilm formation. Sub-inhibitory concentrations of 41 42 the aminoglycoside gentamicin or arsenite significantly decreased the growth rate and yield, but only arsenite caused a significant increase of the lag phase. Hydrogen peroxide 43 44 presented higher disinfection effectiveness against aminoglycoside susceptible than against resistant strains, an effect that was not observed for UV or chlorine. Although this 45 conclusion needs validation based on a larger number of isolates, including clinical, the 46 47 results suggest that aminoglycoside resistance may be associated with traits that influence *Ralstonia* spp. fitness in the environment. 48

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KEYWORDS

52 aminoglycosides resistance; biofilm; disinfectants; kinetics; sub-inhibitory concentrations

56 INTRODUCTION

Members of the phylum Proteobacteria, mainly some families and genera of the classes 57 Alpha-, Beta- and Gammaproteobacteria are amongst the most prevalent bacteria in water 58 59 habitats (Vaz-Moreira et al. 2014; Vaz-Moreira et al. 2017). Some species of the genus 60 Ralstonia within the class Betaproteobacteria, in particular the species Ralstonia pickettii and *R. mannitolilytica*, are frequently observed in aquatic habitats, specifically in 61 wastewater, potable water, surface water and mineral water (Becerra-Castro et al. 2015; 62 63 Falcone-Dias et al. 2012; Ryan et al. 2011; Vaz-Moreira et al. 2017). These Ralstonia species are comprised of ubiquitous bacteria that have been found in a wide variety of 64 environments, such as in plastic (Poly Vinyl Chloride) pipes forming biofilms structures, 65 66 aerospace samples, purified water, saline solutions, skin disinfectants, biological samples, caps of blood culture bottles or even in human patients in the respiratory tract or the oral 67 cavity (Adley et al. 2005; Anderson et al. 1990; Boutros et al. 2002; Coenye et al. 2002; 68 69 Coman et al. 2017; Daxboeck et al. 2005; Koenig and Pierson 1997; Kulakov et al. 2002; Labarca et al. 1999; Maroye et al. 2000; McNeil et al. 1985; Mijnendonckx et al. 2013; 70 71 Riley and Weaver 1975; Ryan et al. 2006; Stelzmueller et al. 2006; Verschraegen et al. 1985). The ubiquitous character of these bacteria is related to the requirement for minimal 72 nutrient resources and explains the transmission from various sources to humans 73 74 (Daxboeck et al. 2005). The capacity to grow in moist environments and to form biofilm has also been proposed as a reason for the *Ralstonia* spp. persistence in some environments 75 (Adley et al. 2005). Motility, a characteristic of *Ralstonia* spp., is also sometimes 76 associated with the increased capacity to form biofilm (Guttenplan and Kearns 2013; 77 O'Toole and Kolter 1998). Although the decrease of motility after prolonged preservation 78

79	and sub-culture has been reported in <i>Ralstonia</i> spp. (Ryan 2009; Vaneechoutte et al. 2001),
80	the association between motility and biofilm formation have been proposed in this genus.
81	In particular, in some Ralstonia spp. aerotaxia was observed to regulate the biofilm
82	formation (Yao and Allen 2007).
83	R. pickettii and R. mannitolilytica are not considered primary pathogens and, hence, are not
84	screened in routine monitoring analyses in hospitals (Coenye et al. 2002; Orme et al. 2015;
85	Waugh et al. 2010). Nevertheless, it has been argued that the low frequency of infection
86	episodes attributed to Ralstonia spp. may be a consequence of misidentifications of these
87	bacteria, being suggested that members of this group may be more widespread, invasive
88	and severe than previously thought (Coman et al. 2017; Daxboeck et al. 2005; Ryan et al.
89	2006). A recent study reported an association between the presence of intestinal Ralstonia
90	pickettii and an augmented glucose intolerance in obesity (Udayappan et al. 2017).
91	Although members of R. pickettii and R. mannitolilytica lack major virulence factors, and
92	rarely are reported as causing infection, members of these species have been considered the
93	most pathogenic species of the genus (Vaneechoutte et al. 2004). Indeed, nosocomial
94	outbreaks attributed to R. pickettii or R. mannitolilytica have been reported regularly over
95	the last 30 years (Coman et al. 2017; De Baere et al. 2001; Fernandez et al. 1996; Khajuria
96	et al. 2014; Labarca et al. 1999; Riley and Weaver 1975; Ryan et al. 2006; Vaneechoutte et
97	al. 2001; Verschraegen et al. 1985).
98	In part, the ubiquitous character and potential to infect humans may be associated with the
99	capability to stand environmental stresses observed in R. pickettii and R. mannitolilytica.
100	For instance, the capacity to survive in hospital disinfectants, as chlorhexidine and
101	ethacridine lactate (acrinol) (Ryan et al. 2006) or to participate in bioremediation processes

102	through the breakdown of xenobiotic compounds was described in <i>R. pickettii</i> (Ryan et al.
103	2007). Also metal tolerance seems to be a relevant property of these bacteria illustrated by
104	the fact that the microbiota enriched from a hospital effluent in copper led to the isolation
105	of monospecies cultures of metal resistant Ralstonia spp., in spite of the complex microbial
106	community of such an effluent (Becerra-Castro et al. 2015). The array of antibiotic
107	resistance phenotypes to gentamicin, chloramphenicol, colistin, tobramycin, polymyxin B,
108	and many others observed in members of these species are probably also related with their
109	ubiquity (Daxboeck et al. 2005; Pan et al. 2011; Ryan et al. 2009; Stelzmueller et al. 2006).
110	Given this background information, the occurrence of <i>R. pickettii</i> and <i>R. mannitolilytica</i>
111	throughout the urban water cycle (wastewater and tap water) and in pristine water sources
112	(mineral water) (Becerra-Castro et al. 2015; Falcone-Dias et al. 2012; Kulakov et al. 2002;
113	Ryan et al. 2006; Vaz-Moreira et al. 2017), raises interest on their ecology. Able to
114	withstand the drinking water treatment, bacteria of these species can reach humans through
115	the water consumption. This work was based on the hypothesis that traits, such as tolerance
116	to metals and antibiotics, to disinfection or the capacity to produce biofilm in the presence
117	of antibiotics, may differ among Ralstonia spp. strains and hence affect their response to
118	stress and environmental fitness. As part of the experimental design, a set of five Ralstonia
119	spp. strains, isolated from hospital wastewater, mineral water, and tap water, were tested for
120	their response under the abovementioned stress types.

122 MATERIALS AND METHODS

123 Bacterial strains

Five *Ralstonia* spp. isolates were selected for this study, four *Ralstonia pickettii*, two from 124 125 hospital wastewater, one from mineral water and one from tap water, and one *Ralstonia* 126 *mannitolilytica* from tap water (Table 1). Cultures were maintained and preserved in nutritive (Luria-Bertani) broth supplemented with 15% (v/v) glycerol. 127 128 The identification of the strains was made based on the 16S rRNA gene sequence analysis using the primers 27F and 1492R as previously described (Ferreira da Silva et al. 2007). 129 130 The sequences were compared with the public database EzBioCloud (Yoon et al. 2017). 131 The five strains were characterized based on selected biochemical tests using the 132 commercial kits API 20E, API 20NE, and API ZYM (bioMérieux) following the 133 manufacturer's instructions. Capsule presence was tested by negative staining (McKinney 134 1953) in cultures grown in the absence and presence of sub-inhibitory concentrations of gentamicin. These additional characterizations were done in an attempt to find some traits 135 that could be associated with the antibiotic and metals resistance phenotypes. 136 137 The 16S rRNA gene sequences of the studied strains were compared with other good 138 quality (>1000 bp) sequences of R. pickettii and R. mannitolilytica strains of different origins available in the GenBank (http://www.ncbi.nlm.nih.gov/). The nucleotide sequence 139 analysis was performed using the MEGA6 software (Tamura et al. 2013), based on the 140 141 model of Jukes and Cantor (Jukes and Cantor 1969), and a dendrogram was created using 142 the neighbour-joining method. The iTol software v3.2.4 (Letunic and Bork 2016) was used 143 to represent the isolates source in the dendrogram.

144 Determination of antibiotic and metal resistance phenotypes

145 The antibiotic resistance phenotypes were determined by disk diffusion method as

recommended by the Clinical Laboratory Standards Institute (CLSI 2015), for 12

- 147 antibiotics: nalidixic acid (NA, 30 μg); ciprofloxacin (CIP, 5 μg); streptomycin (STR, 10
- 148 μg); gentamicin (GEN, 10 μg); tetracycline (TET, 30 μg); cephalothin (CP, 30 μg);
- 149 meropenem (MER, 10 μg); ceftazidime (CEF, 30 μg); ticarcillin (TIC, 75 μg); colistin
- sulphate (CT, 50 μg); sulfamethoxazole (SUL, 25 μg) and sulfamethoxazole/trimethoprim
- 151 (SXT, 23.75/1.25 μg). The interpretation criteria (R, resistance; S, susceptible) based on
- inhibition zone diameters were as follows (mm): NA30: $R \le 13$, $S \ge 19$; CIP5: $R \le 15$, $S \ge 21$;
- **153** STR10: R≤11, S≥15; GEN10: R≤12, S≥15; TET30: R≤11, S≥15; CP30: R≤14, S≥18;
- 154 MER10: R≤15, S≥19; CEF30: R≤14, S≥18; TIC75: R≤15, S≥24; CT50: R≤10, S≥11;
- 155 SUL25: $R \le 12$, $S \ge 17$; SXT25: $R \le 10$, $S \ge 16$. In each assay, the reference strain *Pseudomonas*
- aeruginosa DSM 1117 was used for quality control.
- 157 Determination of Minimum Inhibitory Concentrations (MICs)
- 158 The minimum inhibitory concentrations (MICs) were determined using the Etest or the
- 159 microdilution method at 30 °C. The Etest (BioMérieux, France) or MICE (OXOID, United
- 160 Kingdom) were used for the antibiotics gentamicin (CN 256-0.015 μ g/mL, OXOID,
- 161 MA0116F), streptomycin (SM 0.064-1024 µg/mL, BioMérieux, 526800), ceftazidime (TZ
- 162 0.016-256 µg/mL, BioMérieux, 412293), meropenem (MEM 32-0.002 µg/mL, OXOID,
- 163 MA0121F), and sulfamethoxazole (SX $0.064-1024 \mu g/mL$, BioMérieux, 412458). The
- 164 microdilution method was used for tetracycline and metals (Andrews 2001), using bacterial
- suspensions of absorbance 0.08-0.1 at 625 nm in Mueller-Hinton broth supplemented with
- 166 0.1 32 mg/L of tetracycline, 0.001 2 mmol/L of NaAsO₂, 0.01 10 mmol/L of
- 167 NiCl₂· $6H_2O$ or 1 14 mmol/L of CuSO₄· $5H_2O$. For concentrations of CuSO₄· $5H_2O$ above
- 168 5 mmol/L the MICs were tested in Tris-buffered Mueller-Hinton broth. The MICs were

determined as the minimum concentration that inhibited visible bacterial growth after 24 hof incubation.

Based on preliminary distinctive results between the tested strains, the aminoglycoside
gentamicin, and the metal arsenite were selected to assess their effects as stressors and will
be from this point forward designated as stressors. Each of the five strains was assayed in
stressor-free (SF) culture medium and in the presence of gentamicin or arsenite at
concentrations close to the MIC value.

176 Stressors and growth kinetics

177 Cultures were assayed in Mueller-Hinton broth or in this culture medium supplemented 178 with adequate concentrations and volume of stressor solution. Therefore, strains H2Cu2, T6BT1 and L1PA1 were assayed in 125 mg/L gentamicin or 1.1 mmol/L As³⁺; strain 179 H2Cu5 was assayed in 6 mg/L gentamicin or 0.01 mmol/L As³⁺; and strain T6BT10 was 180 181 assayed in 0.4 mg/L gentamicin or 0.01 mmol/L As³⁺. Bacterial suspensions with an initial absorbance of 0.05 at 610 nm (A610) were incubated at 30 °C with orbital shaking (~70 182 rpm) and were monitored every hour until reached the stationary phase (~24 h). Growth 183 curves and kinetic parameters (growth rate, lag phase, and yield) were determined in 184 185 triplicate in independent assays. Growth curves were represented as log values of A610 in function of time. The lag phase was the period of time necessary to start the exponential 186 187 phase. The growth rate (μ) was determined based on the slope of the curve during the exponential growth phase, according to the equation $Ln N_t - Ln N_0 = \mu(t - t_0)$, where N is 188 189 the number of cells at time t. The growth yield corresponded to the maximum A610 reached. 190

The capacity of each strain to form biofilm was tested in modified Luria-Bertani broth 192 193 (mLB) (tryptone 5 g/L, yeast extract 2.5 g/L and sodium chloride 1 g/L) over a range of different stressor concentrations: 0.01, 0.05, 0.5 and 1.1 mmol/L As³⁺; 25, 75 and 125 mg/L 194 of GEN; and 125, 250, 500 and 750 mg/L of STR, concentrations bellow the MICs for the 195 196 strains H2Cu2, L1PA1 and T6BT1; and of 0.01 mmol/L of As³⁺; 0.4 and 6 mg/L of GEN; 197 and 50 mg/L of STR for the strains H2Cu5 and T6BT10. The assays were performed in 198 clear flat bottom 96-well polystyrene microtiter plates (Orange Scientific, Belgium) as 199 described by Simões *et al.* (2007). Briefly, the microtiter wells were filled with 200 μ L of bacterial suspension (A610 = 0.1; prepared from overnight cultures in mLB at 30 °C) in 200 201 mLB or in mLB supplemented with one of the stressors, incubated for 48 h at 30 °C and 202 measured the absorbance at 620 nm (A620) in a microplate reader (FLUOstar optima, BMG Labtech, Germany). After that, the plates were washed with phosphate buffer and air-203 204 dried for 30 min. To assess and compare the biofilm formation, the biomass was fixed with methanol, left to dry, stained with crystal violet, washed again and the dye resuspended 205 206 with glacial acetic acid prior to measuring the absorbance at 570 nm (A570). A negative 207 control consisting of non-inoculated culture medium and a reference culture (Pseudomonas aeruginosa DSM 1117) were included in each assay. Each experiment was performed at 208 209 least six times for each strain. The quantification of the biofilm formation was performed as 210 described by Rode et al. (Rode et al. 2007), through the calculation of a ratio A_{570}/A_{620} , 211 referring to absorbance at 570 nm (to measure the biofilm formation) and absorbance at 620 212 nm (to measure the bacterial growth). The absorbance values were corrected by the subtraction of the respective absorbance measured in the negative control (non-inoculated 213

214	culture medium). With the procedure used, the possible contribution of the growth yield for
215	the capacity to form biofilm was normalized by the use of the ratio A_{570}/A_{620} , referring to
216	absorbance at 570 nm (measure of the biofilm formation) and absorbance at 620 nm
217	(measure of the bacterial growth).
218	Disinfectants and inactivation
219	The effectiveness of the germicide UV radiation, chlorine or hydrogen peroxide, was tested
220	in saline solution (0.85% (w/v) NaCl) bacterial suspensions of $A610 = 0.1$. Suspensions

were prepared from 24 h Plate Count Agar (PCA) cultures. Samples collected at the

beginning and over the assay were cultivated for enumeration on PCA and incubated at 30

°C for 24-48 h.

For UV disinfection was used a germicide UV lamp with a wavelength of 254 nm, under

which were exposed PCA plates onto which were spread $100 \,\mu$ L of a bacterial suspension

with about 10 to 300 CFU/mL. Exposure times were of periods of 0, 15, 30, 45, 60, 90 and

227 150 seconds.

228 To test the effect of chlorine was used a solution of 10 mg/L sodium hypochlorite prepared

from commercial bleach with a concentration of sodium hypochlorite equivalent to 5% (50

230 g/L). Bacterial suspensions (A610 = 0.1) prepared in saline solution were exposed to

sodium hypochlorite at a final concentration of 5 mg/L. A solution of 1.5% (w/v) sodium

thiosulfate was used to neutralize the effect of chlorine at different exposure times of: 0, 2,

233 7, 12, 17, 25 and 60 minutes. Cultures were plated immediately after the addition of the

aneutralizing agent.

The effect of hydrogen peroxide was tested using a 0.1% solution prepared from a 30%
stock (Carlo Erba Reagents, Italy). Bacterial suspensions (A610 = 0.1) were exposed to
hydrogen peroxide at a final concentration of 0.05% (v/v). A freshly prepared solution of
bovine liver catalase (0.1 g/L) was used in a ratio 0.1/5 (v/v) to eliminate residual hydrogen
peroxide (Fiorentino et al. 2015) after exposure times of: 0, 2, 7, 12, 17, 25 and 60 minutes.
Cultures were plated after catalase addition.

241 Statistical analyses

242 The effect of different stressors and the behavior of different strains was compared based 243 on the parametric test one way ANOVA or the nonparametric tests Kruskal-Wallis and 244 Mann-Whitney, depending if the results followed or not a normal distribution. The capacity 245 to form biofilm in the presence and absence of stressors was compared based on the 246 nonparametric test Mann-Whitney. The effect of disinfectants on cells inactivation was 247 compared based on parametric one way ANOVA test with post-hoc test Tuckey. All the statistical analyzes were performed with the SPSS software package, version 23.0 (IBM 248 249 SPSS software, Chicago, IL).

250

251 **RESULTS**

252 *Ralstonia* spp. tolerance to antibiotics and heavy metals

- 253 Based on the 16S rRNA gene sequence analyses, the *R. pickettii* and *R. mannitolilytica*
- strains studied clustered together with others from sources such as plant/animal,
- clinical/human, water and soil, or other environments (e.g. air) (Fig. 1). Although not

256	related to the isolation origin, three phylogenetic subgroups could be distinguished, one that
257	included strains H2Cu2, L1PA1 and T6BT1, sharing a 16S rRNA gene sequence identity of
258	99.7 - 99.9%, other including strain H2Cu5, with a 16S rRNA identity with first group of
259	99.0 - 99.2% and another one of <i>R. mannitolilytica</i> , which, non-surprisingly included the
260	strain T6BT10 with a 16S rRNA gene sequence identity of 97.8 - 98.2% with the <i>R</i> .
261	pickettii isolates tested. These differences were not confirmed at the biochemical phenotype
262	for which the five strains displayed a similar profile (data not shown).
263	All strains were observed to be resistant to colistin and ticarcillin and susceptible to the
264	quinolones (nalidixic acid, ciprofloxacin), sulfonamides (sulfamethoxazole and
265	sulfamethoxazole/trimethoprim), beta-lactams (cephalothin and ceftazidime) and
266	tetracycline. Variable phenotypes were observed for meropenem and aminoglycosides
267	susceptibility. Strains R. pickettii H2Cu5 and R. mannitolilytica T6BT10, both susceptible
268	to gentamicin, differed on the susceptibility to streptomycin observed only in R.
269	mannitolilytica T6BT10. Hence, gentamicin resistance was observed in the group of
270	phylogenetically closely related strains H2Cu2, L1PA1 and T6BT1, with MIC-gentamicin
271	values >256 mg/L, while strains H2Cu5 and T6BT10, in distinct phylogenetic subgroups,
272	presented lower MIC values for both gentamicin and streptomycin (Table 2). MIC values
273	for As ³⁺ were about 30 times higher in gentamicin resistant than in gentamicin susceptible
274	isolates, while no differences among strains were observed for Ni ²⁺ or Cu ²⁺ . This finding
275	suggested that aminoglycoside and arsenite resistance mechanisms might be associated.

276 <u>Stressors and growth kinetic</u>

277	Based on the hypothesis that a common mechanism of resistance could be used by these
278	strains for gentamicin and arsenite, growth kinetic parameters were determined in the
279	absence and in the presence of each of those stressors (Table 3). In the absence of any
280	stressor, the growth rates for the five strains were similar (~0.4 h^{-1}) (Table 3). Either
281	gentamic n or arsenite led to significant ($p < 0.05$) reductions in the growth rate, being the
282	highest reductions observed in the presence of sub-inhibitory concentrations of gentamicin
283	(Table 3). The lag phases in the absence of stressor ranged 0.7-0.9 h. In the presence of
284	arsenite, but not in the presence of gentamicin, these values significantly $(p < 0.05)$
285	increased (to 2.4-3.4 h) in the strains with highest MIC-As ³⁺ values (Table 3). In absence of
286	stressors, growth yield ranged 2.4-2.9. These values that were significantly ($p < 0.05$)
287	reduced in the presence of arsenite for strain H2Cu2 (to 2.2) or in the presence of
288	gentamicin for strains H2Cu2, H2Cu5, L1PA1, and T6BT1 (to 0.5-0.9). In general, the
289	reduction of growth yield was more pronounced in the presence of gentamicin than of
290	arsenite. These differences in the growth parameters in the presence of gentamicin or
291	arsenite suggest that even if a common resistance mechanism is used to grow in the
292	presence of each of those stressors, probably distinct functions are targeted in the cell by
293	the antibiotic or the metal.

294 Effect of stressors in the capacity of biofilm formation

295 The capacity to form biofilm may be an advantage in *Ralstonia* spp. to face adverse

conditions (Adley et al. 2005; Anderson et al. 1990; Di Domenico et al. 2016; Ryan et al.

- 2011). Hence, it was hypothesized that the stressors aminoglycosides and arsenite could
- stimulate the capacity to produce biofilm (Fig. 2). The low concentrations of stressor
- tolerated by the aminoglycoside susceptible strains H2Cu5 and T6BT10 were not observed

300 to induce in those strains an increased capacity to form biofilm. In contrast, the strains 301 resistant to the aminoglycosides gentamicin and streptomycin (H2Cu2, T6BT1 and L1PA1) presented significant increases in the capacity to form biofilm, with increases of 2-4 times 302 for the lower concentrations and 5-11 times for the highest concentrations of 303 304 aminoglycosides tested, in comparison with the non-stressor assays (Fig. 2). In the same way, 1.1 mmol/L arsenite a significant, although lower (1.5-1.9 times), increase of biofilm 305 306 formation. The capacity to form biofilm can be associated with the production of 307 polysaccharide capsules that facilitate the adherence to surfaces and the formation of 308 biofilms (Moxon and Kroll 1990). It was thus hypothesized that the increased capacity to 309 form biofilm could be due to an observable overproduction of capsule polysaccharides in the presence of sub-inhibitory concentration of gentamicin. However, this hypothesis was 310 311 not proved, eventually because the method used to observe capsules was not sufficiently 312 sensitive.

313 Disinfectants and inactivation

314 The hypothesis beyond these assays was that aminoglycoside and arsenite resistant strains 315 would present a higher resilience against the different types of disinfectant - UV radiation, chlorine, and peroxide disinfection (Fig. 3). However, it was observed that only peroxide 316 317 disinfection supported that hypothesis. Neither UV radiation nor chlorine were observed to 318 produce a distinct effect on the gentamicin resistant or susceptible strains (Fig. 3). The UV radiation promoted a reduction of 1-2 log at each 15 min of exposure till the maximum 319 320 period tested of 45 min (Fig. 3A). In the presence of 5 mg/L chlorine it was observed a sharp culture inactivation (2 min), to reach after 7 min of exposure, counts < 10 CFU/mL 321 322 (Log 1) (Fig. 3B). In contrast to the other two disinfectants, hydrogen peroxide revealed

higher antibacterial effectiveness against the gentamicin and arsenite susceptible strains
than against the resistant. Susceptible strains decreased to counts below the quantification
limit (one log-unit) after 7 min of exposure, in contrast to the resistant strains that required
12 min to reach < one log-unit (Fig. 3C).

327

328 **DISCUSSION**

329 *Ralstonia* spp. are ubiquitous, mainly in aquatic environments, including drinking water 330 (Vaz-Moreira et al. 2017), and have been reported as contaminants of clinical sterile 331 solutions or materials (Boutros et al. 2002; Labarca et al. 1999; McNeil et al. 1985) or as 332 the prevalent species in hemodialysis water samples (Vincenti et al. 2014). Ralstonia 333 *pickettii* strains from different clinical and environmental origins were reported as being 334 resistant to gentamicin, ticarcillin and meropenem, although highly susceptible to 335 ciprofloxacin, tetracycline and sulfamethoxazole/trimethoprim (Ryan and Adley 2013), 336 confirming the phenotypes of the strains examined in this study. The observation that phylogenetically and phenotypically close Ralstonia strains, isolated from distinct aquatic 337 338 environments, differed on the susceptibility to gentamicin (Table 1 and 2), suggested that it 339 might be due to gene acquisition. Indeed, in a parallel study, whole genome sequencing from strains H2Cu2 and H2Cu5 showed that only the aminoglycoside resistant isolate 340 341 contains genes associated with resistance to arsenic, and toxic compounds, encoding 342 lysozyme inhibitors, or phages/prophages receptors (Vaz-Moreira et al. 2016). 343 The correlation observed between the resistance to arsenite and gentamicin, may also be an 344 indication of a possible mechanism of co-resistance (genetic linkage between two or more

345 resistance genes) or cross-resistance (same genetic determinant confers resistance to both 346 antibiotics and metals), frequently reported for heavy metals and antibiotics (Baker-Austin 347 et al. 2006; Dib et al. 2008; Seiler and Berendonk 2012; Zhou et al. 2015) specially in contaminated environments (Ahemad and Malik 2013). 348 Supposedly the physiological response to arsenite and aminoglycosides involves diverse 349 mechanisms, as the response for biofilm formation and growth kinetic did not respond in 350 351 the same mode for both antimicrobials (Table 2). However, the stress imposed by sub-352 inhibitory concentrations of aminoglycosides or of metals increased the capacity to form 353 biofilms of the aminoglycoside resistant isolates (Fig. 2). This can be due to an unspecific 354 stress response, not related with the resistance mechanism, in particular enhanced production of extracellular polymeric substances (EPS) which lead to cell adhesion (Baker-355 356 Austin et al. 2006; Donlan 2002; Donlan and Costerton 2002; Lindsay and von Holy 2006), and biofilm formation (Balaban 2008; Donlan and Costerton 2002; Lindsay and von Holy 357 2006). Similar results were observed for *Pseudomonas aeruginosa* and *Escherichia coli* 358 isolates, increasing their capacity to form biofilm in the presence of aminoglycosides (Aka 359 and Haji 2015; Hoffman et al. 2005). However, Paul et al. (2014) observed a negative 360 influence of 0.01 mmol/L arsenite on the capacity to form biofilm of *Pseudomonas* spp. or 361 362 *Rhizobium* spp. strains. Some authors refer to the importance of the bacteria motility on the 363 capacity to form biofilm (Guttenplan and Kearns 2013; O'Toole and Kolter 1998; Yao and 364 Allen 2007). Indeed, *Ralstonia* spp. are motile and this property might influence the 365 variable capacity to form biofilm observed in the present study. However, it was not possible to assess differences in motility capabilities in the tested strains or to investigate if 366 stressors interfere with flagella and therefor with biofilm formation. Considering the history 367

368 of *Ralstonia* spp. as important colonizers of highly oligotrophic environments, other369 conditions to explore could be the effect of nutritional stress.

370 In contrast to what was observed for hydrogen peroxide, the survival to UV radiation or

371 chlorine disinfection did not differ in aminoglycoside resistant or susceptible strains (Fig.

372 3). This observation is probably related with the inactivation mechanisms involved,

373 DNA/RNA damage for UV, cell metabolism for chlorine and oxidative stress for hydrogen

peroxide (Estrela et al. 2002; Hijnen et al. 2006; McDonnell and Russell 1999). These

375 results suggest that the mechanisms involved in the aminoglycoside resistance are probably

not directly associated with the mechanisms of survival to UV radiation or chlorination.

377 Studies performed with a higher number of isolates, from different origins may give more

378 consistency to these results. The capability of *Ralstonia* spp. to survive in some

379 environments subjected to disinfection processes, as for example the wastewater or

drinking water systems, is probably determinant for their capacity to spread or be

transmitted to humans. It is curious to note that aminoglycoside resistant strains have

increased capacity to form biofilm in the presence of some environmental stressors, since

this may be a relevant factor to facilitate the survival and spread of aminoglycoside

384 resistant *Ralstonia* spp. strains in the environments subjected to stress conditions, as

antimicrobial challenges. These characteristics combined with the oligotrophic character

may contribute for the *Ralstonia* ubiquity in aquatic habitats.

387

388 CONCLUSIONS

389 The aminoglycosides resistance was associated with the highest tolerance to arsenite.

390	Sub-inhibitory concentrations of gentamicin or arsenite significantly decreased the growth
391	rate and yield, while arsenite but not gentamicin caused a significant increase of the lag
392	phase. The biofilm formation was stimulated in the presence of aminoglycosides or
393	arsenite, in the aminoglycoside resistant but not in the susceptible strains.
394	Disinfection with UV or chlorine presented identical effectiveness in aminoglycoside
395	resistant or susceptible strains. In contrast, hydrogen peroxide presented higher
396	effectiveness against aminoglycoside susceptible than resistant strains.
397	The results support the hypothesis that antibiotic resistance is associated with improved
398	tolerance to stress.
399	
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TABLES:

581	Table 1.	Group	of isolates	tested in	this study	
					2	

				Abundance in the			
Strain	Species	Source of	Isolation medium and conditions	source (order of	Reference		
		isolation		magnitude, CFU's/mL)			
		Hospital	Culture enrichment in modified Luria-	103	D. C. () () () () ()		
H2Cu2	R. pickettu	wastewater	Bertani broth with Cu ²⁺ (2.5 mmol/L)	103	Becerra-Castro <i>et al.</i> 2015		
H 2C ₂ 5	P. piakattii	Hospital	Culture enrichment in modified Luria-	103	Pagama Castro et al 2015		
H2Cu5	к. ріскенн	wastewater	Bertani broth with Cu^{2+} (2.5 mmol/L)	10-	Becenta-Castro et al. 2013		
T 1DA 1		Min anal matan	Pseudomonas isolation agar with	10	Falaana Diag at al 2012		
LIPAI	к. ріскетні	32mg/L amoxicillin		10.	Faicone-Dias <i>et al.</i> 2012		
T6BT1	R. pickettii	Tap water	Tergitol 7-agar	10-1	Vaz-Moreira et al. 2013		
T6BT10	R. mannitolilytica	Tap water	Tergitol 7-agar	10-1	Vaz-Moreira et al. 2013		

583 Table 2. Minimum Inhibitory Concentrations (MICs) for antibiotics and metals determined

	MICs									
Strain	GEN	STR	ТЕТ	MER	CEF	SUL	As ³⁺	Ni ²⁺	Cu ²⁺	
	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mmol/L)	(mmol/L)	(mmol/L)	
H2Cu2	> 256	>1024	1	>32	6	24	1.4	4	12	
H2Cu5	6	56	0.25	6	6	4	0.05	4	12	
L1PA1	> 256	>1024	1	>32	6	24	1.4	4	12	
T6BT1	> 256	>1024	1	16	8	24	1.4	4	12	
T6BT10	0.5	4	8	>32	4	4	0.05	4	12	

584 for the *Ralstonia* spp. strains under study.

585 GEN, gentamicin; TET, tetracycline; MER, meropenem; CEF, ceftazidime; SUL, sulfamethoxazole; STR,

586 streptomycin and metal salts of As^{3+} , Ni^{2+} and Cu^{2+}

Table 3. Variations on the bacterial growth parameters growth rate, phase lag and yield, under sub-inhibitory concentrations of arsenite

Strain	Growth rate (per hour)					Phase Lag (hours)				Yield (A610)								
concentration)	SF		As^{3+}		GEN		SF		As ³⁺		GEN		SF		As^{3+}		GEN	
H2Cu2 125 mg/L GEN or 1.1 mmol/L As ³⁺	0.4±0.03	1;a,b	0.3±0.02	2;a	0.1±0.01	3;a	0.9±0.2	1;a	3.4±0.5	2;a	2.0±0.7	1,2;a	2.9±0.1	1;a	2.2±0.2	2;a	0.5±0.1	3;a
H2Cu5 6 mg/L GEN or 0.01 mmol/L As ³⁺	0.3±0.01	1;a	0.3±0.04	1;a	0.1±0.01	2;b	0.7±0.4	1;a	1.0±0.3	1;b	2.2±2.5	1;a	2.5±0.3	1;a	2.0±0.3	1;a	0.9±0.2	2;a
L1PA1 125 mg/L GEN or 1.1 mmol/L As ³⁺	0.4±0.04	1;a,b	0.3±0.04	1;a,b	0.2±0.01	2;c	0.7±0.4	1;a	3.0±0.9	2;a	0.8±0.1	1;a	2.9±0.4	1;a	2.5±0.5	1;a	0.8±0.1	2;a
T6BT1 125 mg/L GEN or 1.1 mmol/L As ³⁺	0.4±0.02	1;a,b	0.3±0.01	2;a	0.2±0.01	3;b	0.9±0.3	1;a	2.4±0.3	2;a	1.4±0.4	1;a	2.7±0.1	1;a	2.4±0.3	1;a	0.7±0.1	2;a
T6BT10 0.4 mg/L GEN or 0.01 mmol/L As ³⁺	0.4±0.01	1;b	0.4±0.01	1,2;b	0.4±0.01	2;d	0.8±0.1	1;a	0.8±0.1	1;b	0.7±0.1	1;a	2.4±0.5	1;a	2.1±0.4	1;a	1.9±0.6	1;b

588	(As^{3+}) or gentamicin	(GEN) or con	trol conditions	(stressor free, SF).
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589 A610, bacterial suspension absorbance at 610 nm;

590 Statistically significant differences between stress conditions (SF, As^{3+} and GEN) are indicated by the numbers: 1, 2, 3; and significant differences between

591 strains are indicated by the letters: a, b, c, d.

FIGURES:

- **Fig. 1** Environmental distribution of *Ralstonia pickettii* and *Ralstonia mannitolilytica*,
- 596 including the strains studied (in red).

599	Fig. 2 Influence of stressors on the ability of biofilm formation, for the strains H2Cu2 (A),
600	T6BT1 (B) and L1PA1 (C). The quantification of the biofilm formation was performed
601	through the calculation of a ratio A_{570}/A_{620} , referring to absorbance at 570 nm (measure of
602	the biofilm formation) and absorbance at 620 nm (measure of the bacterial growth). The
603	non-inoculated control presented a ratio A_{570}/A_{620} of 1.0±0.1; and the <i>P. aeruginosa</i>
604	presented ratios A ₅₇₀ /A ₆₂₀ of: 3.1±0.7 for stressor-free (SF); 2.2±0.9 for gentamicin (GEN)
605	6 mg/L; 2.8±0.7 for streptomycin (STR) 50 mg/L; 0.2±0.09 for meropenem 4 mg/L;
606	1.9 ± 0.9 for arsenite (As) 0.01 mmol/L; 0.3 ± 0.05 for copper 6.0 mmol/L; and 0.7 ± 0.2 for
607	nickel 2.5 mmol/L.
608	

Fig. 3 Bacterial inactivation with A) UV radiation, B) chlorine (5 mg/L), and C) hydrogen
peroxide (0.05%).