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Hunting for cultivable *Micromonospora* strains in soils of the Atacama Desert

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15 Abstract (150-250)

Innovative procedures were used to selectively isolate small numbers of 16 17 Micromonospora strains from extreme hyper-arid and high altitude Atacama Desert soils. Micromonosporae were recognized on isolation plates by their ability to produce 18 19 filamentous microcolonies that were strongly attached to the agar. Most of the isolates 20 formed characteristic orange colonies that lacked aerial hyphae and turned black on 21 spore formation whereas those from the high altitude soil were dry, blue-green and covered by white aerial hyphae. The isolates were assigned to seven multi- and eleven 22 single-membered groups based on BOX-PCR profiles. Representatives of the groups 23 were assigned to either multi-membered clades that also contained marker strains or 24 formed distinct phyletic lines in the Micromonospora 16S rRNA gene tree, many of the 25 isolates were considered to be putatively novel species of *Micromonospora*. Most of the 26 isolates from the high altitude soils showed activity against wild type strains of *Bacillus* 27 subtilis and Pseudomonas fluorescens while those from the rhizosphere of Parastrephia 28 quadrangulares and from the Lomas Bayas hyper-arid soil showed resistance to UV 29 30 radiation.

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Keywords: *Micromonospora*, Atacama Desert, BOX-PCR, polyphasic taxonomy, UV
 radiation.

37 Introduction

38 Given their importance in biotechnology, bioprospecting and ecology, actinobacteria remain a source of interest to the microbiological community (Goodfellow and Fiedler 39 2010; Demain 2014; Barka et al. 2016). Actinobacteria were initially isolated from the 40 Atacama Desert over fifty years ago (Cameron et al. 1966; Opfell and Zebal 1967) 41 though the first extensive survey of these organisms in this desert was only reported 42 recently (Okoro et al. 2009). Okoro and her colleagues isolated small numbers of 43 44 filamentous actinobacteria from arid, hyper-arid and extreme hyper-arid Atacama Desert soils using selective isolation procedures and polyphasic taxonomic methods 45 which showed that a high proportion of the isolates belonged to the genera 46 Amycolatopsis, Lechevaleria and Streptomyces many of which were assigned to 47 putative new species and contained novel non-ribosomal peptide synthase genes. 48 Subsequent studies have shown that Atacama Desert habitats are not only a rich source 49 of novel streptomycetes but also of rare and poorly studied taxa, some of which produce 50 bioactive compounds and have been given validly published names (Bull et al. 2016; 51 Busarakam et al. 2014, 2016; Goodfellow et al. 2017; Idris et al. 2017a; Idris et al. 52 2017b; Trujillo et al. 2017). Streptomyces leewenhoekii strains (Busarakam et al. 2014), 53 for instance, synthesize novel macrolactone and polyketide antibiotics (Nachtigall et al. 54 2011; Rateb et al. 2011a; Rateb et al. 2011b) and chaxapeptin, a new lasso peptide 55 (Elsayed et al. 2015) while the type strain of *Lentzia chajnantorensis* (Idris et al. 2017b) 56 produces novel diene and monoene glycosides, several of which show anti-HIV 57 integrase activity (Wichner et al. 2017). Complementary metagenomic analyses of 58 Atacama Desert habitats have revealed a remarkable actinobacterial diversity most of 59 which has not been detected using culture-dependent methods (Bull et al. 2017; Idris et 60 al. 2017c). Improved selective isolation and cultivation methods are needed to isolate 61 components of this diversity, not least Micromonospora strains which are known to be a 62 rich source of new specialist metabolites and have the potential to defend plants against 63 64 root-infecting fungi (Carro et al. 2018a).

65 Micromonospora (Ørskov 1923), the type genus of the family Micromonosporaceae (Krasil'nikov 1938; Zhi et al. 2009), currently encompasses 79 66 species with validly published names (Parte 2014) many of which have been 67 circumscribed using polyphasic methods (Genilloud 2012; Carro et al. 2018b) though 68 the genus remains underspeciated (Carro et al. 2012a; Carro et al. 2013a). 69 70 Micromonosporae have been isolated from diverse natural habitats (Genilloud 2012), notably from rhizosphere soil (Carro et al. 2013b; Thawai et al. 2016) from desert 71 locations in China (Ding et al .2013) and from tissues of a broad range of plants, such as 72 73 Triticum aestivum (Coombs and Franco 2003), Zea mays (Shen et al. 2014), and Parathelypteris beddomei (Zhao et al. 2017); as well as from nitrogen-fixing root 74 75 nodules of actinorhizal plants (Trujillo et al. 2006; Carro et al. 2013a) and legumes (Trujillo et al. 2007; Garcia et al. 2010; Trujillo et al. 2010; Carro et al. 2018b). 76

It is apparent from culture-independent studies that micromonosporae form a small, but integral part of actinobacterial communities in Atacama Desert habitats (Bull et al. 2017; Idris et al. 2017c). The failure to isolate them from such habitats may reflect the use of isolation media mainly designed to be selective for streptomycetes (Busarakam 2014; Idris 2016). The primary aims of the present study were to isolate
micromonosporae from diverse Atacama Desert soils using procedures designed to be
selective for members of this taxon and to determine whether any such isolates
represented putatively novel taxa.

85

86 Materials and Methods

87 Sampling sites

Soil samples (Table 1) were collected from several locations in the Atacama Desert (Fig. 1) between 2010 and 2016 (ATB, MG) and an additional one in 2012 (Professor Luis Cáceres, University of Antofagasta). The samples were collected aseptically using spatulas sterilized in the field with ethanol and contained in sterile polycarbonate bottles. Following transport to the UK the samples were stored at 4°C. Further details of the sampling sites can be found elsewhere (Bull et al. 2017; Idris et al. 2017c).

94 *Selective isolation*

Three isolation procedures were used: a) Phenol-heat protocol (P-H): a gram of each of 95 the soil samples was diluted in 0.85% sodium chloride solution containing 1.5% phenol 96 (Hayakawa et al. 1991), the preparations shaken on a rotary shaker for an hour, 97 incubated at 70°C for 40 minutes in a water bath (Nonomura and Ohara, 1969), shaken 98 again at room temperature for 2 hours, serial dilutions prepared in the saline solution, 99 and the resultant preparations shaken at room temperature prior to spreading over chitin-100 vitamin (CHV) (Zhang 2011), Gause number 1 (G nº 1) (Gause et al. 1983), humic acid-101 vitamin (HV) (Havakawa and Nonomura 1987), M65 (DSMZ medium number 65), 102 R2A (Reasoner et al. 1979) and Zhang' starch soil extract agar (ZSSE) (Zhang 2011), all 103 of the media were supplemented with 50µg/ml of cycloheximide and 50µg/ml of 104 nalidixic acid; b) sprinkle protocol (S): 0.5g of soil particles of individual samples 105 which had been preheated at 120°C for 15 minutes were sprinkled directly over the 106 107 isolation media and c) a standard dilution plate protocol (D): dilutions of each soil sample in saline were used to inoculate each of the isolation media following the 108 procedure described by Goodfellow et al. (1967). All of the inoculated plates were 109 incubated at 28°C and examined weekly for up to 28 days for the presence of 110 Micromonospora colonies that were detected, using a stereoscopic microscope, and then 111 transferred to the medium from which they were isolated but without the antibiotics, the 112 resultant plates were incubated at 28°C for 15 days. The 45 isolated strains were 113 maintained on either M65 or ZSSE agar and in 20% v/v glycerol at -80°C for long-term 114 preservation. 115

116 *Dereplication of isolates*

Genomic DNA for BOX-PCR was extracted from all the isolates using a Bacterial DNA 117 Extraction Kit (Sigma) following the instructions of the manufacturer and BOX-PCR 118 fingerprint profiles generated using modifications of the methods described by 119 Versalovic et al. (1994) and Trujillo et al. (2010). To this end, Bioline 2x MiFiTM mix 120 was used for PCR amplification in a final volume of 25 µl per reaction following the 121 manufacturer's recommendations; the thermal cycling parameters were: 7 minutes at 122 95 °C, 30 cycles of 1 minute at 94 °C, 1 minute at 52 °C and 3 minutes at 72 °C followed 123 by a final extension at 72 °C for 10 minutes. Five microliters of each PCR product was 124

loaded onto a 2% agarose gel containing 10 µl of GelRedTM (Crisafuli et al. 2015) per
100 ml and electrophoresis run at 70 V for 3 hours in freshly prepared 1x TBE-EDTA
buffer at pH 8.0 using a Bio-Rad Pac 300 power supply; a DNA molecular weight
marker 1kb HyperLadderTM (Bioline) was used as the molecular size standard. After
electrophoresis, gels were photographed, stored on disk as TIFF files, and manually
aligned into 9 multi- and 10 single-membered similarity groups.

131 Phylogeny

132 PCR-mediated amplification of genomic DNA of 20 representatives of the BOX-PCR groups was conducted using the universal primers 27F and 1522R (Lane 1991). Bioline 133 2x MiFiTM mix was used for PCR amplification in a final volume of 50 µl per reaction 134 following the manufacturer's recommendations. The thermal cycling parameters were: 135 9 minutes at 95 °C, 30 cycles of 1 minute at 94 °C, 1 minute at 56 °C and 2 minutes at 136 72 °C followed by a final extension at 72 °C for 10 minutes. The PCR products were 137 purified using a QIAquick[®] PCR purification kit, according to the manufacturer's 138 instructions (Qiagen) and sequenced using the EZ-seq Barcode Service (Macrogen). 139 The resultant sequences (around 1400 nucleotides) were manually aligned using 140 MUSCLE (Edgar 2004) and then compared with corresponding sequences of 141 Micromonospora type strains retrieved from the EzBioCloud server (Yoon et al. 2016). 142 143 Phylogenetic distances were calculated with the Kimura 2-parameter model (Kimura 1980) and tree topologies inferred using the neighbour-joining (Saitou and Nei 1987) 144 and maximum-likelihood algorithms (Felsenstein 1981) and one thousand bootstrap 145 replications. Tree reconstructions were carried out using MEGA 7 software (Kumar et 146 147 al. 2016).

148 Stress tests

149 All of the strains were examined for their ability to grow in the presence of various 150 concentrations of sodium chloride (1, 3, 5, 7 and 9%, w/v) and at a range of temperature (4, 10, 20, 28, 37 and 45 °C) and pH values (4.5, 5.5, 6.5, 8.0 and 9.0) using M65 and 151 ZSSE agar as basal media; pH values were determined using phosphate buffers, as 152 described previously (Carro et al. 2012b). All of the plates, apart from those from the 153 temperature tests, were incubated at 28°C for 15 days. The strains were also examined 154 for their ability to grow following exposure to 100 mJoules for 30 and for 60 minutes in 155 a UV chamber (Biorad) using *Geodermatophilus poikilotrophi* DSM 44209^T as the 156 positive control (Montero-Calasanz et al. 2014)) set at a wavelength of 254 nm (UVC). 157 The capacity of the strains to grow under anaerobic conditions was tested on M65 and 158 ZSSE agar plates that were incubated for 28°C for 10 days in anaerobic atmosphere 159 generation bags (Sigma-Aldrich 68061). 160

161 *Antibiotic sensitivity assays*

The isolates were examined for their ability to inhibit the growth of wild type strains of 162 Bacillus subtilis, Escherichia coli and Pseudomonas fluorescens using a standard plug 163 assay (Fiedler, 2004). Lawns of each of the isolates were prepared on yeast extract-malt 164 extract and oatmeal agar (International Streptomyces Project [ISP] media 2 and 3; 165 Shirling and Gottlieb 1966) plates incubated at 28°C for 14 days. Agar plugs taken from 166 the incubated plates were transferred to sterile square Petri Dishes and overnight Luria 167 broth cultures [OD₆₀₀ of approximately 0.6] of each of the wild type strains in molten 168 nutrient agar added to a final concentration of OD₆₀₀ of 0.0125. The inoculated dishes 169

170 were incubated at 28°C and examined for zones of inhibition around the agar plugs after

171 24 and 48 hours.

172 Results and Discussion

173 Isolation of micromonosporae

The ability to isolate specific fractions of actinobacterial communities from the 174 Atacama Desert biome is of paramount importance especially since metagenomic 175 analyses have revealed a staggering degree of actinobacterial diversity therein: the vast 176 majority of which has gone undetected using culture-dependent procedures (Idris et al. 177 2017c), this partly reflects the use of isolation media that favour the growth of 178 streptomycetes (Williams et al. 1984). Isolation procedures designed to recover specific 179 actinobacterial genera from environmental samples are many and varied but tend to 180 reflect the biological properties of the target organisms (Goodfellow 2010), as was the 181 case in the present study. It is, for instance, known that Micromonospora spores are 182 activated by heat pretreatment regimes, are resistant to phenol and germinate on nutrient 183 rich media such as humic acid-vitamin agar (Hoskisson et al. 2000; Shen et al. 2014; 184 Carro et al. 2018b). It is, therefore, encouraging that over half of the Micromonospora 185 strains were isolated from the soil samples using the phenol-heat procedure, albeit on 186 187 several isolation media (Table 2).

188 Micromonosporae growing on the isolation plates were recognized, under the stereoscopic microscope, by their ability to produce filamentous microcolonies that 189 were strongly attached to agar, these colonies were readily distinguished from those of 190 191 aerobic, endospore-forming bacilli (data not shown). It was particularly interesting that most of the Micromonospora strains came from the ALMA 4 sample as complementary 192 193 culture-independent studies have shown that soil samples from this location contain micromonosporal propagules (Bull et al., 2017). The strains isolated from this site 194 proved to be unusual as they formed dry, filamentous blue-green colonies covered by 195 white aerial hyphae following growth for three weeks on M65 and ZSSE media (Fig. 1). 196 The ability of these isolates to form aerial hyphae is interesting as some 197 198 Micromonospora type strains have been shown to contain putative homologous genes to those associated with aerial hyphae formation and spore maturation in streptomycetes 199 (Carro et al. 2018a). Strains isolated from soil samples taken from the three remaining 200 locations showed a typical micromonosporal phenotype, one characterized by the 201 202 production of filamentous orange colonies that lack aerial hyphae and turn blue-black upon spore formation (Genilloud 2012). Five of these strains were isolated from 203 204 extreme hyper-arid soil collected from the Lomas Bayas region (Idris et al. 2017c), one 205 of the driest areas in the Atacama Desert; all of these isolates were recovered by plating serial dilutions of the soil onto M65 agar. In turn, the 13 strains isolated from the 206 207 rhizosphere of Parastrephia quadrangularis (Compositae, tribe Asteraceae) were obtained either by sprinkling mineral particles or spreading serial dilutions of the 208 209 phenol-heat protocol onto HV and ZSSE agar and incubating for four weeks at 28°C. Micromonospora strains were recovered using all three selective isolation procedures 210 211 (Table 2).

212 BOX-PCR profiles

The BOX-PCR profiles of the isolates encompass considerable genetic diversity with 213 fragments ranging from 0.05 to 2.0 kb (Fig. 2) thereby providing further evidence that 214 this method is effective in distinguishing between *Micromonospora* strains (Maldonado 215 et al. 2008; Trujillo et al. 2010). The isolates were assigned to seven multi- and 11 216 single-membered groups defined at the 60% similarity level (Table 3). Five of the 217 multi-membered groups and five of the singletons were composed of strains isolated 218 from the ALMA 4 soil samples, the largest of these taxa, group XI, contained seven 219 isolates. In turn, the strains isolated from P. quadrangularis rhizosphere soil were 220 assigned to a multi-membered group that encompassed 10 strains and to two single 221 membered groups; the five Lomas Bayas strains formed one multi- and 3 single-222 membered taxa while the single isolates from ALMA 5 and Yungay Core Region 223 samples gave unique profiles (Fig. 2). It is interesting that the multi-membered groups 224 225 only contained strains isolated from a single location and that few strains were clones (Fig. 2; Table 3). 226

227 *Phylogenetic analyses*

Almost complete 16S rRNA gene sequences (1372 - 1507 nucleotides [nt]) were 228 generated for the 20 isolates chosen to represent the BOX-PCR groups (Table 3). The 229 generic assignment of all of these strains was confirmed as they were recovered in the 230 Micromonospora 16S rRNA gene tree (Fig. 3). The 16S rRNA gene sequence 231 similarities between these isolates and the type strains of their closest phylogenetic 232 neighbours fell within the range 98.4 to 99.9 % (Table 3). Several of the isolates were 233 assigned to well-supported clusters in the Micromonospora tree though none of these 234 taxa included strains isolated from more than one location. Ten out of the twelve 235 ALMA 4 strains formed a well-supported phyletic line within a weakly supported clade 236 237 that encompassed the type strains of *Micromonospora costi* (Thawai 2015), their nearest 238 neighbour, and Micromonospora fulviviridis (Kroppenstedt et al. 2005); it is interesting that the isolates of this clade were recovered from different samples of ALMA 4 soil 239 240 using two of the three isolation protocols and several selective isolation media (Table 241 2). Isolate 5R2A7, one of the remaining ALMA strains, formed a well-supported clade 242 together with the type strain of Micromonospora coriariae (Trujillo et al. 2006), its nearest phylogenetic neighbour, and with Micromonospora cremea (Carro et al. 2012b) 243 244 whereas isolate ATA32, the remaining ALMA 4 strain, formed an unsupported clade 245 with the type strain of Micromonospora narathiwatensis (Thawai et al. 2007) though it was most closely related to Micromonospora eburnea (Thawai et al. 2005). 246

It is interesting that two out of the three strains recovered from the rhizosphere 247 of P. quadrangularis, isolates STR1-7 and STR1s-6, were loosely associated with the 248 type strains of Micromonospora lupini (Trujillo et al. 2007), Micromonospora taraxaci 249 (Zhao et al. 2014) and Micromonospora violae (Zhang et al. 2014) which were isolated 250 from a root nodule of Lupinus angustifolius and the roots of Taraxacum mongolicum 251 and Viola philippica, respectively (Fig. 3). Moreover, the final strain, isolate STR1s-5, 252 was most closely related to the type strain of Micromonospora ureilytica (Carro et al. 253 2016b), which was isolated from a root nodule of *Pisum sativum*. In turn, two of the 254 four strains isolated from the extreme hyper-arid Lomas Bayas soil, isolates LB19 and 255 LB32, were phylogenetically close to the type strain of Micromonospora saelicesensis 256 (Trujillo et al. 2007), an isolate from a root nodule of L. angustifolius. It is also 257 258 interesting that isolate LB39 was recovered in the well supported clade that included the type strains of *M. chokoriensis, M. taraxaci* and *M. violae* (Fig. 3); the final strain from
the Lomas Bayas soil, isolate LB4, formed a well-supported clade with the type strain
of *Micromonospora chalcea* (Foulerton 1905, Orskov, 1923) which was isolated from
air. It can be seen from the Figure 3 that the sole strain from the extreme hyper-arid
Yungay Core soil, isolate Y6-2, forms a well-supported clade with the type strain of *Micromonospora pisi* (Garcia et al. 2010), an isolate from a root nodule of *Pisum*sativum.

266 To date, representatives of dereplicated groups of actinobacteria isolated from Atacama Desert soils which show low pairwise 16S rRNA gene sequence similarities, 267 (<90%; Meier-Kolthoff et al. 2013) with the type strains of their nearest phylogenetic 268 neighbour have been invariably assigned to new species when subject to polyphasic 269 analyses. as exemplified by Lechevalieria, Pseudonocardia 270 taxonomic and Streptomyces species (Okoro et al. 2010; Busarakam et al. 2014; Trujillo et al. 2017). 271 272 This cut-off point has also proved to be a reliable indicator for the presumptive recognition of novel Micromonospora species (Trujillo et al. 2007; Garcia et al. 2010; 273 Carro et al. 2016a) though pairwise 16S rRNA gene sequence similarities very much 274 higher than the 99.0% threshold have been found to be indicative of new 275 Micromonospora species, as exemplified by M. taraxaci and M. violae (Zhang et al. 276 277 2014; Zhao et al. 2014). Given these indicators it seems likely that further comparative taxonomic analyses will show that most, if not all, of the representatives of the BOX-278 PCR groups will be found to represent novel Micromonospora species. Indeed, isolates 279 280 ATA38, ATA39, and ATA43 from ALMA4, isolate STR1-7 from the rhizosphere of P. quadrangularis, and isolate Y6 2 from Yungay are cases in point as they share 16S 281 rRNA gene sequence similarities with their closest phylogenetic neighbours at or below 282 the 99.0% threshold (Table 3). 283

284 Adaptation to extreme conditions

285 In general, the pH and temperature profiles of the isolates were typical of Micromonospora strains (Genilloud 2012) as they grew from 20 to 37°C, at pH 6.0 to 286 8.0, but not at 4, 12 or 45°C or below pH 5.0. None of the isolates grew under anaerobic 287 conditions, but most of them were able to grow in the presence of 1%, w/v sodium 288 289 chloride; six out of the thirteen strains isolated from the rhizosphere of P. 290 quadrangularis grew in the presence of 3%, w/v sodium chloride. It was also interesting that only 30% of the samples from ALMA soil grew on M65 agar following exposure to 291 292 UV light (UVC) at 100 mJoules/second for 30 minutes, this number increased to 60% on ZSSE agar. Similarly, high percentages were observed for strains isolated from the 293 294 rhizosphere and Lomas Bayas soils on M65 (64 and 80%) and ZSSE (50 and 60%) agar, 295 respectively. These results are interesting as it has been shown that the type strain of *Modestobacter caceresii*, an isolate recovered from a soil sample from the Yungay Core 296 Region, has the capacity to protect and repair damaged caused by UV radiation 297 (Busarakam et al. 2016). Interestingly, the sole Micromonospora strain obtained from 298 this area, isolate Y6-2, grew after an hour exposure to UV light at 100 mJoules/second, 299 300 only around 20% of the isolates grew under this condition. Although UVC radiation no 301 longer reaches the Earth's surface it has been used as a selective tool with reference to early life on the planet and high altitude biology. Recently Paulino-Lima et al. (2016) 302 screened desert soils, including Atacama soil, for UVC-resistant bacteria and 303 interestingly over 40% of their isolates were members of the phylum Actinobacteria; 304

Micromonospora strains were not recovered by these authors. Consequently, the distinctive radiation resistance of *Micromonospora* strain Y6-2 suggests that it could be a novel target for detailed radiation-resistant physiological and biochemical researches that should be extended to include the other radiation stressors naturally present on-at the Earth surface. The study of these radiations will be of interest for future studies of the strains and to determine if key osmoprotectant proteins are implicated in the process as previously shown in *Rhodobacter* (Pérez et al 2017).

312 Approximately half of the isolates showed weak activity in the plug assays against the wild type strains of B. subtilis and P. fluorescens though no activity was 313 shown against the E. coli strain. However, it is important that these preliminary 314 315 antibiotic sensitivity studies are extended as the genomes of *Micromonospora* type strains have much greater potential to synthesize novel specialised metabolites than 316 previously realised (Carro et al. 2018a). Indeed, novel strains of Micromonospora and 317 corresponding strains of other genera classified in the family Micromonosporaceae 318 should be given greater prominence in the search for new classes of bioactive 319 compounds, notably antibiotics that are needed to control drug resistant pathogens. 320

It can be concluded that the isolation procedures used in this study provide an effective way of isolating small numbers of putatively novel *Micromonospora* species from Atacama Desert soils. Indeed, innovative selective isolation procedures based on the biological properties of target organisms are needed to cultivate elements of the extensive actinobacterial dark matter detected in soils of the Atacama Desert landscape using culture-independent methods (Bull et al. 2017; Idris et al. 2017c)

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334

335 **Conflict of Interest**

The authors declare that they have no conflict of interest.

- 337
- This article does not contain any studies with human participants or animals performed by any of the authors.
- 340

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566 **Figure legends**

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568 Figure 1. A. Sampling sites: Lomas Bayas (LB), Salar de Tara (ST), Atacama Large Millimeter Array observatory (ALMA) and Yungay (Y6); B. Salar de Tara site; C. 569 Parastrephia. quadrangularis plant growing at the Salar the Tara; D. Typical orange 570 colony of a Micromonospora isolate; E. Typical black Micromonospora colony 571 following spore production; F-H. Morphology observed for some Atacama 572 micromonosporae; F. Sporulating Micromonospora colonies; G. Mountain-shaped 573 574 Micromonospora isolates; H. Colony from ALMA4 isolate covered by white aerial 575 hyphae.

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Figure 2. BOX–PCR fingerprints showing the genetic diversity of *Micromonospora*strains isolated from Atacama Desert soils.

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Figure 3. Neighbour-joining phylogenetic tree based on almost complete 16S rRNA gene sequences showing relationships between the *Micromonospora* strains and between them and *Micromonospora* type strains. Numbers at the nodes indicate the levels of bootstrap support (%), only values above 50% are shown. Asterisks indicate branches of the tree that were also recovered in the maximum-likelihood tree. *Catellatospora citrea* was used as the outgroup. Bar, 0.005 substitutions per nucleotide position.

Sampling site and code	Collection date	Altitude (masl)	Latitude (°S)	Longitude (°W)	Habitat	N° of isolates
ALMA 4 ATA, 4G	20.10.12	4000	23°03'31''	67°52'27''	Subsurface soil (30 cm)	25
ALMA 5 5R2A	20.10.12	5046	23°00'49''	67°45'31''	Surface soil (2 cm)	1
LB 3	24.02.14	1500	23°24'27''	69°31'03''	Extreme hyper-arid surface soil (2 cm)	5
Cerro Aguas Blancas, Y6	13.11.10	1047	24°06'18''	70°01'15''	Extreme hyper-arid subsurface soil (30 cm)	1
STR1	05.10.16	4174	23°03'97''	67°18'87''	Rhizosphere of Parastrephia quadrangularis	13
	site and code ALMA 4 ATA, 4G ALMA 5 5R2A LB 3 Cerro Aguas Blancas, Y6	site and codeCollection dateALMA 420.10.12ATA, 4G20.10.12ALMA 520.10.12SR2A24.02.14LB 324.02.14Cerro Aguas Blancas, Y613.11.10	site and codeCollection dateAltitude (masl)ALMA 4 ATA, 4G20.10.124000ALMA 5 SR2A20.10.125046LB 324.02.141500Cerro Aguas Blancas, Y613.11.101047	site and code Collection date Altitude (masl) Latitude (°S) ALMA 4 20.10.12 4000 23°03'31'' ATA, 4G 20.10.12 5046 23°00'49'' ALMA 5 20.10.12 5046 23°00'49'' LB 3 24.02.14 1500 23°24'27'' Cerro 13.11.10 1047 24°06'18'' Aguas Blancas, Y6 Y6 Y6	site and code Conection date Aintude (masl) Lantude (°S) Longitude (°W) ALMA 4 20.10.12 4000 23°03'31'' 67°52'27'' ATA, 4G 20.10.12 5046 23°00'49'' 67°45'31'' ALMA 5 20.10.12 5046 23°00'49'' 67°45'31'' LB 3 24.02.14 1500 23°24'27'' 69°31'03'' Cerro 13.11.10 1047 24°06'18'' 70°01'15'' Aguas Blancas, Y6 Y6 Y6 Y6	site and codeConlection dateAlfitude (masl)Lantude (°S)Longitude (°W)HabitatALMA 4 ATA, 4G20.10.12400023°03'31''67°52'27'' (°W)Subsurface soil (30 cm)ALMA 5 SR2A20.10.12504623°00'49''67°45'31'' (67°45'31'')Surface soil (2 cm)LB 324.02.14150023°24'27''69°31'03'' (9°31'03'')Extreme hyper-arid surface soil (2 cm)Cerro Aguas Plancas, Y613.11.10104724°06'18'' (70°01'15'')Extreme hyper-arid subsurface soil (30 cm)STR105.10.16417423°03'97'' (67°18'87'')Rhizosphere of Parastrephia

Table 1. Location, sampling sites and dates of collection of soil samples from theAtacama Desert.

+The Atacama Large Millimeter Array (ALMA) Observatory is situated on Cerro
Chajnantor: ALMA is operated as an international partnership which includes the
European Southern Observatory (ESO). Permission to collect soil samples from Cerro

594 Chajnantor was given by the Director of the ESO.

Table 2. Source and substrate mycelial colour of the isolated *Micromonospora* strains.

	Isolates	Isolation protocol	Isolation media	Colour of substrate mycelium
	4G51, 4G53, 4G55, 4G57	D	G nº 1	green
-	ATA32, ATA34, ATA47 ATA48	P-H P-H	ZSSE CHV	orange orange
ALMA 4	ATA31, ATA33, ATA35, ATA36a, ATA36b, ATA37, ATA38, ATA39, ATA40, ATA42, ATA43, ATA44, ATA45	P-H	ZSSE	green
	ATA50	P-H	CHV	green
	ATA46, ATA51	P-H	HV	green
	ATA52	P-H	HV	black
ALMA 5	5R2A7	D	R2A	orange
Lomas Bayas	LB 4, LB 19, LB 32, LB 39, LB 41	D	M65	orange
Yungay core region	Y6_2	P-H	HV	orange

	STR1-41, STR1-74, STR1-85	P-H	HV	orange
Salar de Tara	STR1s-5, STR1s-6, STR1s-7, STR1s-11	S	HV	orange
(rhizosphere)	STR1-7, STR1-71, STR1-72	P-H	ZSSE	orange
	STR1s-13a, STR1s-14, STR1s-16	S	ZSSE	orange
506				

Table 3. Assignment of isolates to BOX groups and determination of their nearestphylogenetic neighbours.

BOX group	Isolates	Representative isolates	% Similarity to closest phylogenetic type strain		
I	ATA31, ATA33, ATA45, ATA47,ATA48, ATA52	ATA 45, ATA52	<i>M. costi</i> 99.1		
11	ATA35, ATA37, ATA40	ATA40	M. costi	99.1	
111	ATA38	ATA38	M. costi	98.4	
IV	ATA39	ATA39	M. costi	98.4	
v	ATA42	ATA42	M. costi	99.1	
VI	ATA43	ATA43	M. terminaliae	98.8	
VII	ATA32, ATA34	ATA32	M. eburnea	99.4	
VIII	LB4, LB41	LB4	M. chalcea	99.6	
IX	LB19	LB19	M. saelicesensis	99.8	
х	LB32	LB32	M. saelicesensis	99.8	
XI	LB39	LB39	M. chokoriensis	99.9	
XII	ATA36, ATA44, 4G51, ATA 46, 4G55, 4G53, 4G57	4G51, 4G57	M. costi	99.1	
XIII	ATA51a, ATA51b	ATA51b	M. costi	99.1	
XIV	5R2A7	5R2A7	M. coriariae	99.8	
xv	STR1-7, STR1-41, STR1- 72, STR1-74, STR1-85, STR1s-7, STR1s-11, STR1s-13A, STR1s-14, STR1s-16	STR1-7	M. chokoriensis	99	
XVI	STR1s-5	STR1s-5	M. chokoriensis	99.4	
XVII	STR1s-6	STR1s-6	M. ureilytica	99.6	
XVIII	Y6-2	Y6-2	M. pisi	98.8	