Metabolomic comparison using *Streptomyces* spp. as a factory of secondary metabolites

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Abstract: Understanding extremophiles and their usefulness in biotechnology involves studying 16 their habitat, physiology and biochemical adaptations, as well as their ability to produce biocata-17 lysts, in environments that are still poorly explored. In northwestern Peru, which has saline lagoons 18 of marine origin Pacific Ocean, the other site is from the coast of Brazil of the Atlantic Ocean. Both 19 environments are considered extreme. The objective of the present work was to compare two dif-20 ferent strains isolated from these extreme environments at the metabolic level using molecular net-21 work methodology through the Global Natural Products Molecular Social Network (GNPS). In our 22 study, the MS/MS spectra from the network were compared with GNPS spectral libraries, where 23 the metabolites were annotated. Differences were observed in the molecular network presented in 24 the two strains of Streptomyces spp. coming from these two different environments. Within the an-25 notated compounds from marine bacteria, the metabolites characterized for Streptomyces sp. B-81 26 from Peruvian marshes were lobophorins A (1) and H (2), as well as divergolides A (3), B (4) and C 27 (5). Streptomyces sp. 796.1 produced different compounds, such as glucopiericidin A (6) and dehy-28 dro-piericidin A1a (7). The search for new metabolites in underexplored environments may there-29 fore reveal new metabolites with potential application in different areas of biotechnology. 30

Keywords: Genome mining; marine environments; molecular networking; bacterial extremophiles; 31 secondary metabolites 32

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1. Introduction

Approximately 22,500 biologically active substances are obtained from microorgan-35 isms, 45% of which are represented by Actinomycetes and 70% of which are Streptomyces 36 metabolites; however, information on substances isolated from microorganisms inhabit-37 ing saline environments is scarce [1,2,3]. The search for these microorganisms has been 38 mainly associated with the production of antibiotics and antitumor substances [4]. Halo-39 philic and halotolerant strains show heterogeneous physiological characteristics for dif-40 ferent genera because these bacteria can synthesize secondary metabolites to cope with 41 the high salinity and extreme temperature conditions of their environments [5,6]. These 42 extreme conditions favor the development of metabolic competitiveness for the produc-43 tion of enzymes, providing adaptation to the high salinity of the environment [4]. The 44 production of bioactive molecules from actinobacteria other than Streptomyces allows ex-45 tending the search for new molecules and interactions. During the last five years, constant 46

Citation: Flores Clavo, R.; Kelyene Pereira, A.; Ruiz Quiñones, N; Henrique Costa, J.; Pacheco Fill, T.; Fantinatti Garboggini, F. Metabolomic comparison using *Streptomyces* spp as a factory of secondary metabolites.*xxxxxxxxx, 20*, x. https://doi.org/10.3390/xxxxx

Academic Editor: Firstname Lastname

Received: date Accepted: date Published: date

Article

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Copyright: © 2022 by the authors. Submitted for possible open access publication under the terms and conditions of the Creative Commons Attribution (CC BY) license (https://creativecommons.org/license s/by/4.0/). isolation efforts of microorganisms from various environmental niches have resulted in
the recovery of thousands of members of the actinobacteria affiliated with different taxonomic classes [5]. The members of the genus *Streptomyces* stand out as producers of the
largest amount of secondary metabolites already described [6]. Various studies have revealed an approach combining LC–MS/MS and molecular networking as a rapid analytical method for the identification of compounds [7]; this microbial group has a unique ability to produce new products, mainly antibiotics [8,9].

[8] Investigated the metabolic profiling of a cultured endophyte strain (Streptomyces 54 sp. HKI0576) by HPLC–MS and revealed a complex metabolome, Streptomyces sp. 12A35 55 was isolated from deep sea sediment collected from the South China Sea and showed 56 promising antibacterial activities attributed to the spirorotronate antibiotics lobophorins 57 B, F, H, and I from a marine origin [9]; Piericidins and Glucopiericidins A and B isolated 58 from the culture broth of Streptomyces pactum S48727 [10], a new natural product of the 59 lobophorin family designated lobophorin K, from cultures of the marine actinobacteria 60 Streptomyces sp. M-207 [11]; A cryptic BGC for type I polyketides was activated by meta-61 bolic engineering methods, enabling the discovery of a known compound, lobophorin 62 CR4, of deep-sea-derived Streptomyces olivaceus SCSIO T05 [12]. 63

A study reported in Salar de Huasco in the Atacama Desert, considered a polyex-64 treme environment in Chile, [13] described that species of the genus Streptomyces were 65 dominant, and a preliminary study [14] present in the salt ponds of Bayovar and Morrope 66 in Peru showed physicochemical characteristics common to extremophilic bacteria and 67 ample potential to produce bioactive compounds. These extreme conditions encourage 68 the development of metabolic competitiveness for the production, for example, of en-69 zymes [15]. The application of various targeted studies on the biosynthesis of natural 70 products allows a better understanding of the full potential of the producing microorgan-71 isms, thus increasing the return on investment in the search for new compounds [16]. 72

In this study, we used a molecular networking approach for the rapid detection of 73 molecules from two strains of *Streptomyces* spp. obtained from extreme environments: Saloons northwestern of Perú and Cabo Frio in Brazil (Figure 1). The detection method, 75 based on UHPLC–MS/MS combined with data analysis using Global Natural Product Social Molecular Networking (GNPS), led to the annotation of very different compounds 77 produced by each of the *Streptomyces* spp. 78



Figure 1. The geographical distribution of *Streptomyces* spp. location with a description of growth80in solid medium. (1) Sediment of a Bayovar saline lagoon located in northwestern Peru and (2) star-81fish in Cabo Frio (RJ) in Brazil.82

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2. Results and Discussion

2.1. Phylogenetic analysis of 16S RNAr from two Streptomyces spp.

The 16S rRNA sequences of the two marine Streptomyces species analyzed in this 86 study, Streptomyces sp. B-81 collected in 2015 in Peru and Streptomyces sp. 796.1 collected 87 in Brazil 2012, were determined and compared by MEGA11.0 software. The phylogenetic 88 tree obtained from the two Streptomyces strains was mainly divided into two characteristic 89 branches. The samples mentioned before were constructed against 29 Streptomyces strains 90 according to EZBioCloud and are presented in Figure 2. The strain obtained from the Pe-91 ruvian saline lagoon was determined to be Streptomyces sp. B-81 (MW562807), whose phy-92 logenetically related sequence identity (1345 bp 92.7%, gene sequence identity) groups to 93 the type strain *S. olivaceus* NRRL B-3009^T with 99,93%, *S. pactum* NBRC 13433^T and boot-94 strap value 97% (Fig. 2 and Table S1 and Table S2). However, for the strain isolated from 95 a marine invertebrate in Brazil, Streptomyces sp. 796.1 (MG654686) sequence identity (1370 96 bp 93.4%, gene sequence identity similarity) is phylogenetically related to the type strains 97 S. buecherae AC541^T with 99,70%, S. youssoufiensis X4^T and S. zagrosensis HM 1154^T with 98 99,48%, and Streptomyces philanthi triangulum^T (DQ3752) with 99,45% gene sequence iden-99 tity similarity and a bootstrap value of 85% (Figure 2 and Tables S1 and S2). Subsequently, 100 each part was further subdivided, and most strains were well separated by clustering un-101 der each species, generating indications that they may be new species. Existing aquatic 102 ecosystems are relatively unexplored but have been reported to harbor a great diversity 103 of microorganisms, which may be reflected in their broad biosynthetic potential [17]. 104

Streptomyces violaceoruber NBRC 12826^T (AB184174) Streptomyces tricolor NBRC 15461^T (AB184687) Streptomyces anthocianicus NBRC 14892^T (AB184631) 96 Streptomyces rubrogriseus LMG 20318^T (AJ781373) Streptomyces lienomycini LMG 20091^T (AJ781353) Streptomyces violaceorubidus LMG 20319^T (AJ781374) Streptomyces tendae ATCC 19812^T (D63873) 97 97 97 Streptomyces tritolerans DAS 165^T (DQ345779) Streptomyces marokkonensis Ap1^T (AJ965470) 97 Streptomyces thinghirensis DSM 41919^T (FM202482) 97 Streptomyces malachitospinus NBRC 101004^T (AB249954) Streptomyces hyderabadensis OU-40^T (FM998652) 97 Streptomyces sp. B-81 (MW562807) 90 97 Streptomyces olivaceus NRRL B-3009^T (JOFH01000101) 96 Streptomyces pactum NBRC 13433^T (AB184398) Streptomyces parvulus NBRC 13194^T (AB184326) Streptomyces lomondensis NBRC 15426^T (AB184673) Streptomyces nigra 452^T (MG572975) Streptomyces coeruleorubidus ISP 5145^T (AJ306622) 97 Streptomyces albogriseolus NRRL B-1305^T (AJ494865) 94^L Streptomyces muensis MBRL 179^T (JN560155) 96 Streptomyces spinoverrucosus NBRC 14228^T (AB184578) 97 Streptomyces abikoensis NBRC 13860^T (AB184537) Streptomyces luteireticuli NBRC 13422^T (AB249969) 96 97 Streptomyces varsoviensis NRRL ISP-5346^T (JOBF01000056) 96 Streptomyces zagrosensis HM 1154^T (JF917242) Streptomyces iranensis HM 35^T (FJ472862) 97 Streptomyces buecherae AC541^T (CP060404) Streptomyces youssoufiensis X4^T (FN421338) 9 - Streptomyces philanthi triangulum^T (DQ375802) 93 ⁸⁵ *Streptomyces* sp. 796.1 (MG654686) Pseudomonas hunanensis LVT (JX545210)

0.02

Figure 2. Phylogenetic tree of Streptomyces spp. strains B-81 and 796.1. The evolutionary history 126 was inferred using the neighbor-joining method [18]. The percentage of replicate trees in which 127 the associated taxa clustered together in the bootstrap test (1000 replicates) are shown next to 128 the branches. The tree is drawn to scale, with branch lengths similar to evolutionary distances 129 used to infer the phylogenetic tree. The evolutionary distances were computed using the Ki-130 mura 2-parameter method [19] and are in units of the number of base substitutions per site. 131 The analysis involved 32 nucleotide sequences. All positions containing gaps and missing data 132 were eliminated. There were a total of 1498 positions in the final dataset. Evolutionary analyses 133 were conducted in MEGA11 [20]. Bar, 0.02 substitutions per nucleotide, Pseudomonas hunanen-134 sis LV^T (JX545210) was used as the outgroup. 135

2.2. MS/MS-Based Molecular Networks of Streptomyces spp.

The metabolites of the crude extracts obtained from each of the *Streptomyces* spp. 137 samples were prepared and analyzed by UHPLC–MS/MS, and the resulting data 138 were analyzed through multivariate data analyses to compare both strains using LC– 139 MS (Fig. 3). The data showed a clear separation between both *Streptomyces* spp. 140

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strains. In particular, using principal component analysis (PCA), Streptomyces sp. B-141 81 was separated from *Streptomyces* sp. 796.1 by PC1 and PC3, explaining ~ 40% of 142 the variance (Fig. 3A). Another multivariate analysis, partial least squares-discrimi-143 nant analysis (PLS-DA), showed clear differentiation between both Streptomyces 144 strains using components 1 and 2 (Fig. 3B), and chemotaxonomic classification based 145 on Streptomyces using liquid chromatography-electrospray ionization-tandem mass 146 spectrometry (LC-ESI-MS/MS) combined with multivariate statistical analysis 147 demonstrated that metabolite-based chemotaxonomic classification is an effective 148 tool for distinguishing Streptomyces spp. and for determining their species-specific 149 metabolites [21]. 150

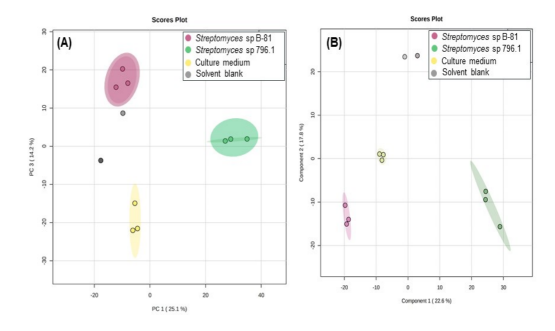


Figure 3. Multivariate data analysis of LC–MS/MS: (A). Principal component analysis (PCA)151and (B). partial least square-discriminant analysis (PLS-DA) comparing both *Streptomyces*152strains. *Streptomyces* sp. B81 is in pink, *Streptomyces* sp. 796.1, in green, the culture medium153R2A, in yellow and the solvent blank is in white.154

Mass spectrometric data obtained from these analyses were then used to gener-155 ate molecular networking coupled to in silico tools (Networking annotation propaga-156 tion (NAP) (https://ccms-ucsd.github.io/GNPSDocumentation/nap/) and MolNetEn-157 hancer (https://ccms-ucsd.github.io/GNPSDocumentation/molnetenhancer/). Data 158 obtained from these analyses were then used to generate more detailed molecular 159 networking (Fig. 4A). Our analyses have presented several chemical classes, such as 160 lipids and lipid-like molecules, organic nitrogen organoheterocyclic compounds, or-161 ganic acids, organosulfur compounds, polyketides, and benzenoid compounds (Fig. 162 4B). 163

Microorganisms in the environment can produce a wide range of secondary metabolites that are natural products with diverse chemical structures and that will perform a variety of functions acting as antibiotics, antitumour agents, cholesterol-lowering agents, etc. [22]. According to the LC–MS/MS analysis, the number of ions can be classified according to the strain source. In all 1005 detected ions, ~ 30% are only from *Streptomyces* sp. B-81 and ~ 64,5% from *Streptomyces* sp. 796.1 (Fig. 4C). Thus, compounds of the same chemical class are grouped in the same cluster, 170

6 of 19

demonstrating a broad biomolecule profile for Streptomyces sp. B-81 (pink), including 171 the production of lobophorins A (1) and H (2), annotated based on their accurate 172 masses, with specific masses of 1157.6379 (1) and 753.4320 (2), respectively; metabo-173 lites were annotated as hits in the GNPS database or manually by accurate mass anal-174 yses, which showed mass errors below 0.10 ppm (Table 1). The cluster of the GNPS 175 database indicated the production of lobophorin A by Streptomyces sp. B-81 176 (MW562807) strain, and fragmentation profile with typical fragments at 97.06, 183.11, 177 517.29 m/z (Fig. S1), the metabolite lobophorin H presented the precursor at m/z178 753.4320 (Fig S2), and compounds that belong to the divergolide family, including 179 divergolides A (3), B (4) and C (5), based on their precursor ions with m/z values of 180 554.2750 (3), 536.2650 (4), and 550.2440 (5), respectively. The fragmentation profile 181 with typical fragments at divergolide C showed the main fragments m/z 182.08, 182 332.22, 398.32 (S3 – S5 Figs). In a previous study of Bayovar's saloons *Streptomyces* sp. 183 B-81 (MW562807) demonstrated the most promising activity, and six biomolecules 184 (Cholic Acid, Lobophorin A, B, E, K and six compounds of type Furano) [14]. The 185 MS/MS spectra of the network metabolites were compared with the GNPS spectral 186 libraries and identified for Streptomyces sp. 796.1, two substances were annotated as 187 glucopiericidin A (6) and dehydropiericidin A1a (7), with precursor ions at m/z188 578.333 (7) and 414.26 (8), which were identified based on their fragmentation pro-189 files with typical fragments at *m*/*z* 161.13, 182.09, 398.26 and 135.11, 182.09, 330.20, 190 respectively (Figs S6-S7) (Fig. 4D and Table 1). In a study carried out on the marine 191 sediment of Saint Peter and Saint Paul's archipelago in Brazil, dereplication analysis 192 of the extracts showed the presence of several compounds identified as piericidins A 193 and C as well as glucopiericidin A, produced by Streptomyces sp. BRA-199 [23] 194

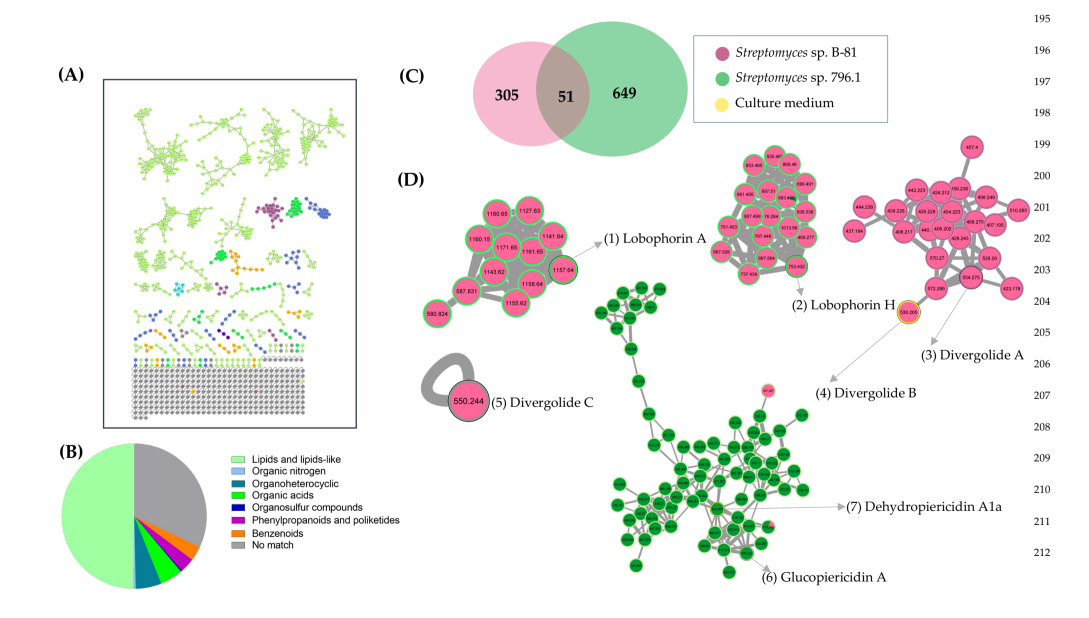


Figure 4. (A) MolNetEnhancer of Streptomyces strains LC-MS/MS data. (B) Pie chart based on213the chemical classes found on the LC-MS/MS data. (C) Euler diagram based on the ion distri-214bution of both strains. (D) Classical molecular networking of both Streptomyces strains.215

Table 1. Mass error (0,10 ppm) of observed and calculated m/z values of known compounds found in clusters based on216MS/MS data obtained from UHPL-HRMS.217

Ν	GNPS NEW-ID	Compound	Molecular Formula [M+H]+	m/z theorethical	<i>m</i> / <i>z</i> observed	error (ppm)	Strain
1	5653	Lobophorin A	C61H93N2O19	1157.6367	1157.6379	1.032275	B-81
2	5178	Lobophorin H	C42H61N2O10	753.4320	753.43200	0.096890	B-81
3	4097	Divergolide A	C31H40NO8	554.2748	554.27500	0.281449	B-81
4	3981	Divergolide B	C31H38NO7	536.2642	536.26500	1.344486	B-81
5	4068	Divergolide C	C31H36NO8	550.2435	550.24400	0.828724	B-81
6	4214	Glucopiericidin A	C31H48NO9	578.3323	578.3330	1.108359	796.1
7	3145	Dehydropiericidin A1a	C25H36NO4	414.263885	414.2650	2.691521	796.1

Error (0,10 ppm) = observed m/z – calculated m/z.

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Ding and colleagues in 2018 took an approach combining LC-MS/MS and mo-219 lecular networks as a rapid analytical method to detect six compounds. Aplysiatoxins 220 from marine cyanobacteria were described with potential new analogs. Another 221 study of MS/MS data obtained from fractions on the public GNPS platform suc-222 ceeded in dereplicating new glycolipopeptides called characellides, cyanocobalamin 223 and poecillastrins of a deep-sea tetractinellid sponge, Characella pachastrelloides 224 [24,25]. Initial chemical investigation of the extract obtained from *Streptomyces* spp. 225 indicated the production of compounds 1-7. Each node within the clusters represents 226 an ionized metabolite, and dereplication was performed by comparison with the m/z227 values of known lobophorins, divergolides and piericidin-related analogs (Table 2). 228 Lobophorins A and B were discovered by Jiang et al. (1999) [26] to Belize onboard 229 the Columbus research ship. Lobophorin C and D were isolated from a new actino-230 mycete strain named # CNC-837, identified as Streptomyces carnosus strain AZS17, 231 isolated from the coastal waters of the East China Sea [27]. Two polyketides, spiro-232 tetronate classified as lobophorins H and I, were isolated from *Streptomyces* sp. 233 1053U.I.1a.3b [28], as well as the other substances of the Lobophorin family (Table 2). 234 The family of compounds named Divergolides isolated from the stem of the man-235 grove tree Streptomyces sp. HKI0576 grown by HPLC-MS revealed a complex metab-236 olome [8]. Four ansamycins, named divergolides T–W, isolated from the mangrove-237 derived actinomycete Streptomyces sp. KFD18 [29], a substance named 3'-deoxytalop-238 iericidin A1 or glucopiericidin A, was isolated from *Streptomyces* sp. soil sample col-239 lected in Gotemba City, Shizuoka, Japan [30], 13-hydroxyglucopiericidin A and the 240 analog Glucopiericidin A were isolated from the fermentation broth of Streptomyces 241 sp. OM-5689 [31] (Table 2). 242

Table 2. A list of known compound families related to *Streptomyces* spp. with compound bioactivity with *m/z* val

ues.

Compound	Molecular Formula [M+H]+	Mass Molecular	Mass calculated	Origins	Reference
Lobophorin A	C61H93N2O19	1157.636705	1157.6379	Streptomyces sp. CNC-837	B-81, [26]
Lobophorin B	C61H91N2O21	1187.610884	1187.6114	Streptomyces sp. CNC-837	[26], [14]
Lobophorin C	C61H90N2O21	1186.603059	1209.5958	Streptomyces sp. AZS 17	[27]
Lobophorin D	C61H93N2O19	1157.636705	1157.6379	Streptomyces sp. AZS 17	[27]
Lobophorin E	C61H91N2O20	1171.615970	1171.6165	<i>Streptomyces</i> sp. SCSIO 01127	[32]

Journal xxxxx 2022, 20, x FOR PEER REVIEW

Lobophorin F	C54H78N2O17	1026.529500	1049.5198	<i>Streptomyces</i> sp SCSIO 01127	[32]
Lobophorin G	C63H94N2O20	1198.639445	1199.6472	Streptomyces sp. MS100061	[33]
Lobophorin H	C42H61N2O10	753.432073	753.43200	<i>Streptomyces</i> sp. 1053U.I.1a.3b	B-81, [28]
Lobophorin I	C61H89N2O21	1185.596209	1185.5958	<i>Streptomyces</i> sp. 1053U.I.1a.3b	[28]
Lobophorin K	C61H93N2O20	1173.631620	1173,6322	Streptomyces sp. M-207	[34]
Lobophorin CR1	C61H92NO20	1144.617647	1180.6032	<i>Streptomyces</i> sp. 7790_N4	[35]
Lobophorin CR2	C61H90N2O22	1202.597974	1247.5702	Streptomyces sp. 7790_N4	[35]
Lobophorin CR3	C61H90N2O23	1218.592889	1241.5832	Streptomyces sp. 7790_N4	[35]
Lobophorin CR4	C52H77N2O15	969.531846	-	Streptomyces olivaceus SCSIO T05	[36]
Lobophorin L	C54H80N2O16	1013.5618	1013.5581	Streptomyces sp. 4506	[37]
Lobophorin M	C48H70N2O13	882.487242	883.4951	Streptomyces sp. 4506	[37]
Lobophorin N	C61H92N2O18	1141.64336	1141.6418	<i>Streptomyces</i> sp. HDN1844000	[38]
Divergolide A	C31H39NO8	554.274844	554.27500	Streptomyces sp. HKI0576	B-81, [8]
Divergolide B	C31H37NO7	536.264279	536.26500	Streptomyces sp. HKI0576	B-81, [8]
Divergolide C	C31H33NO8	550.243544	550.24400	Streptomyces sp. HKI0576	B-81, [8]
Divergolide D	C31H35NO8	535.232645	535.2326	Streptomyces sp. HKI0576	[8]
Divergolide T	C31H37NO7	536.264300	536.2641	Streptomyces sp. KFD18	[29]
Divergolide U	C31H37NO8	550.244600	550.2438	Streptomyces sp. KFD18	[29]
Divergolide V	C31H37NO7	536.264000	536.2640	Streptomyces sp. KFD18	[29]
Divergolide W	C31H37NO7	534.249000	534.2490	Streptomyces sp. KFD18	[29]
Piericidin A-A1	C25H37-39NO4	401.268636	401.2686	Streptomyces sp. 16-22	[39]
Piericidin B-B1	C26H39NO4	415.284286	-	Streptomyces mobaraensis	[40]
Piericidins B1N-oxide	C26H39NO5	431.279201	446.2897	Streptomyces sp. MJ288-OF3	[41]
Piericidin C7	C28H41NO5	457.294851	472.3062	Streptomyces sp. YM14-060	[42]
Piericidin C8	C29H43NO5	471.310501	486.3223	Streptomyces sp. YM14-060	[42]
Glucopiericidin A	C31H48NO9	578.332359	578.3330	Streptomyces spp.	796.1, [30]
Dehydropiericidin A	C25H36NO4	414.263885	414.2650	Streptomyces spp.	796.1, [31]
13- hydroxyglucopiericidin A	C31H47NO10	579.316374	593.3163	Streptomyces sp. OM-5689	[43]
Glucopiericidin C	C30H45NO8	533.310895	533.3108	Streptomyces sp. B811	[44]
JBIR-02	C27H39NO3	411.289372	426.2979	Streptomyces sp. ML55	[45]

3. Materials and Methods

3.1. Biological Material

The microorganisms used in this study were preserved by deep freezing at -247 80°C in 30% glycerol in the research collection of the Microbial Resources Division, 248 Pluridisciplinary Center for Chemical, Biological and Agricultural Research 249 (CPQBA) State University of Campinas (UNICAMP), Campinas, SP, Brazil. 250 (https://www.cpqba.unicamp.br/) and collection of CIICAM Research Center 251 (www.ciicam.com). Two bacteria belonging to the genus Streptomyces were identified 252 as Streptomyces sp. 796.1 (MG654686), isolated from a starfish in Cabo Frio (RJ) in 253 Brazil in 2012, Streptomyces sp. B-81 (MW562807) was isolated from the sediment of a 254 Bayovar saline lagoon located in northwestern Peru in 2015. 255

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10 of 19

The isolates were reactivated using R2A medium (KASVI) with artificial sea-256 water (ASW) (MgCl2-6H2O 10.83 g. L⁻¹, CaCl2-2H2O 1.51 g. L⁻¹, SrCl2-6H2O 0.02 g. L⁻¹, 257 NaCl 23.93 g. L⁻¹, Na₂SO4 4.01 g. L⁻¹, KCl 0.68 g. L⁻¹, NaHCO₃ 0.20 g. L⁻¹, KBr 0.098 g. 258 L-1, H₃BO₃ 0.03 g. L-1) with 5% NaCl and incubated at 28°C for 15 days. After incuba-259 tion, the colonies verified for purity were collected from the plate and used for the 260 extraction of genomic DNA. 261

3.2. DNA isolation and PCR amplification of bacteria

The extraction of genomic DNA was performed using the protocol established 263 by Van Soolingen et al. (1991) adapted to the conditions of the laboratory 264 (https://dx.doi.org/10.17504/protocols.io.bsj4ncqw). First, quantification of genomic 265 DNA was performed by comparing the λ DNA at different concentrations in a 1% 266 TBE 1X agarose gel submitted to electrophoresis for 20 min at 5 V per cm². Then, the 267 DNA was kept at -20°C until the amplification of the 16S rRNA gene was performed 268 by polymerase chain reaction (PCR). The specifications of this protocol have been previously described (dx.doi.org/10.17504/protocols.io.brrmm546). 270

3.3. Sequencing and Phylogenetic Analysis of 16

The samples were purified using a minicolumns GFX PCR DNA & gel band pu-272 rification kit (GE Healthcare Bio-Sciences AB Uppsala, Sweden) according to a pre-273 viously described protocol (https://dx.doi.org/10.17504/protocols.io.brzpm75n). The 274 samples were sequenced using an ABI3500XL Series automatic sequencer (Applied 275 Biosystems Foster City, California, USA) in the Laboratory of the Division of Micro-276 bial Resources, Chemical, Biological and Agricultural Pluridisciplinary Research 277 Center (CPQBA), University of Campinas (UNICAMP), Paulínia, São Paulo, Brazil. 278 Sequencing reactions were performed using the Big Dye Terminator Cycle Sequenc-279 ing Ready Reaction Kit (Applied Biosystems) according to the manufacturer's modi-280 fied by the author's protocol (https://dx.doi.org/10.17504/protocols.io.brzpm75n). 281

Partial sequences of the 16S ribosomal RNA gene obtained from each isolate 282 were assembled into a contig using BioEdit 7.0. The sequences of organisms were 283 added to the EZBioCloud 16S Database (https://www.ezbiocloud.net/) using the 284 "Identify" service, and species assignment was based on the closest hit 16S rRNA 285 gene sequences retrieved from the database and related to the unknown organism 286 gene were selected for alignment in the Clustal W program. Phylogenetic analyses 287 were performed using the MEGA version 11.0 program, and the evolutionary dis-288 tance matrix was calculated using the Kimura-2 model parameters. Then, the phylo-289 genetic tree was constructed from the evolutionary distances calculated by the neigh-290 bor-joining method with bootstrap values based on 1000 resamples. 291

3.4. Production of Crude Extracts of Streptomyces spp

The production of crude ethyl acetate extracts of bacterial isolates was per-293 formed in culture media R2A broth (Himedia ref. 1687) for Streptomyces spp. B-81 and 294 796.1 previously published strains as described in а protocol 295 (dx.doi.org/10.17504/protocols.io.br2cm8aw). Streptomyces sp. metabolites were ex-296 tracted with 1000 µL of methanol in an ultrasonic bath for 40 minutes, dried under 297 inert conditions and analyzed by ultrahigh pressure liquid chromatography-mass 298 spectrometry (UHPLC-MS) in a Thermo Scientific QExactive® Hybrid Quadrupole-299 Orbitrap Mass Spectrometer. 300

3.5. LC–MS/MS data acquisition

Analyses were performed in the positive mode with a range of m/z 133-2000, a 302 capillary voltage of 3.4 kV, inlet capillary temperature of 280°C, and S-lens 100 V. A 303 Thermo Scientific column Accucore C18 2.6 µm (2.1 mm x 100 mm) was used as the 304

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stationary phase, and 5 μ L of the sample was injected. The mobile phase was composed of 0.1% formic acid and acetonitrile, and the eluent profile followed a molar ratio of 95/5 up to 2/98 within 10 min, held for 5 min, up to 95/5 within 1.2 min, and held for 8.8 min. The total run time was 25 min for each run, and the flow rate was 0.2 mL x min-1. The data were initially processed with Xcalibur software (version 309 3.0.63) developed by Thermo Fisher Scientific. 310

3.6. LC–MS/MS data analysis and metabolite annotation

Raw data, blanks, and media controls were converted to.mzML using MSCon-312 vert software (http://proteowizard.sourceforge.net). Next, a molecular network for 313 Streptomyces sp. strain metabolites was created using the online workflow 314 (https://ccms-ucsd.github.io/GNPSDocumentation/networking/) on the Global Nat-315 ural Products Social Molecular Networking (GNPS) website platform 316 (http://gnps.ucsd.edu) using the Classical Molecular Networking (CMN) tool. For 317 network creation, the data were filtered by removing all MS/MS fragment ions within 318 +/- 17 Da of the precursor m/z. MS/MS spectra were filtered by choosing only the top 319 6 fragment ions in the +/- 50 Da window throughout the spectrum. The precursor ion 320 mass tolerance was set to 0.02 Da and an MS/MS fragment ion tolerance of 0.02 Da. 321 A network was then created where edges were filtered with a cosine score above 0.65 322 and more than six (6) matched peaks. 323

Furthermore, edges between two nodes were kept in the network if and only if 324 each of the nodes appeared in each other's respective top 10 most similar nodes. Fi-325 nally, the maximum size of a molecular family was set to 100, and the lowest-scoring 326 edges were removed from molecular families until the molecular family size was be-327 low this threshold. The spectra in the network were then searched against GNPS' 328 spectral libraries. The library spectra were filtered in the same manner as the input 329 data. All matches kept between network spectra and library spectra were required to 330 have a score above 0.65 and at least six (6) matched peaks [46]. The resulting molec-331 ular networking is available at: 332

(https://gnps.ucsd.edu/ProteoSAFe/sta-

tus.jsp?task=b7a29b5a78864223bde01fb8af0c5eff). To perform the multivariate data334analysis, the LC-MS data were processed into MZmine online using GNPS Dash-335board (https://ccms-ucsd.github.io/GNPSDocumentation/lcms-dashboard/). The job336is available at:337

https://gnps.ucsd.edu/ProteoSAFe/sta- 338 tus.jsp?task=88a21c1aabb04a1287348300b711ebfd. The quantification table was submitted to the MetaboAnalyst 5.0 platform (https://www.metaboanalyst.ca/) for statistical analysis. 341

The nodes (MS/MS spectra) originating from R2A culture media and blanks 342 (methanol) were filtered from the original network to enable visualization of metabolites. Finally, the final spectral network (.cys) was uploaded in Cytoscape 3.8 to obtain better visualization and editing. 345

4. Conclusions

In this study, we sought to classify two *Streptomyces* species based on their sec-347 ondary metabolites. The dendrograms based on metabolites and 16S rRNA exhibited 348 the same patterns of species discrimination and may be new species. The multivariate 349 data analyses to compare both strains using LC-MS showed a clear separation be-350 tween both *Streptomyces* spp. The MS/MS spectra of the network metabolites were 351 compared with the GNPS spectral libraries, and differences in the molecular network 352 presented in Streptomyces sp. B-81 isolated from the Bayovar saline lagoon located in 353 northwestern Peru were lobophorins A (1) and H (2), as well as divergolides A (3), B 354 Journal xxxxx 2022, 20, x FOR PEER REVIEW

(4) and C (5); *Streptomyces* sp. 796.1 isolated in Cabo Frio (RJ) in Brazil dereplicated 355 two compounds, glucopiericidin A (6) and dehydropiericidin A1a (7). 356

Supplementary Materials: The following supporting information can be downloaded at
www.mdpi.com/xxx/s1. Phylogenetic data of *Streptomyces* sp. B-81 Accession number
(MW562807), and *Streptomyces* sp. 796.1 Accession number (MG654686); MS/MS match be-
tween GNPS database lobophorin A (1), in the GNPS database or manually by accurate deter-
mination of which ion precursor lobophorin H (2) and ions precursor divergolides A (3), B (4)
and C (5) from *Streptomyces* sp. B-81 extract and MS/MS match between GNPS database with
Glucopiericidin A (6) and Dehydropiericidin A1a (7).357

Author Contributions: R.F.C. performed the extraction of Streptomyces sp. B-81 sample, the364isolation, purification, and structure elucidation of compounds, and constructed the molecular365networks A.K.P, wrote and paper; N.R.Q. performed the isolation and production of Strepto-366myces sp. 796.1 extracts. A.K.P. and J.H.C performed the UHPLC–MS/MS analysis and Thermo367Scientific QExactive® Hybrid Quadrupole-Orbitrap Mass Spectrometer and constructed the368molecular networks; T.P.F and F.F.G edited and vetted the paper. All authors read and corrected the paper.369

Funding: This study has been financed by the Concytec-World Bank Project "Improvement371and Expansion of the Services of the National Science Technology and Technological Innova-
tion System" 8682-PE, through its executing unit ProCiencia [contract number 190-2018] and
The postgraduate programs in Genetics and Molecular Biology and CNPQ.371374

Acknowledgments: This work was supported by the Biotechnology Department of the Center375for Research and Innovation in Multidisciplinary Active Sciences -CIICAM and Pluridiscipli-376nary Center for Chemical, Biological and Agricultural Research (CPQBA), State University of377Campinas (UNICAMP), Paulínia, SP, Brazil.378

Conflicts of Interest: The authors declare no conflict of interest. The funders had no role in the379design of the study; in the collection, analyses, or interpretation of data; in the writing of the380manuscript; or in the decision to publish the results.381

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Journal xxxxx 2022, 20, x FOR PEER REVIEW

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SUPPLEMENTARY DATA

Table S1. Complete 16S rRNA sequences of two *Streptomyces* spp.

Strain	No. of	BLASTn closest homolog	Completennes	Variation
	bp	(accession #) organism	/Identity (%)	radio
Streptomyces sp.	<u>1345</u>	Streptomyces olivaceus NRRL B-3009 ^T	100.0/(99.93)	1/1345
B-81		<u>Streptomyces pactum NBRC 13433</u> ^T	99.6/(99.93)	1/1345
		<u>Streptomyces parvulus NBRC 13193⁺</u>	99.2/(99.03)	13/1340
Streptomyces sp.	1370	<u>Streptomyces buecherae</u> AC541 ^T	100/(99.70)	4/1352
796.1		<u>Streptomyces youssoufiensis</u> X4 ^T	100/(99.48)	7/1352
		<u>Streptomyces zagrosensis</u> HM 1154 ⁺	100/(99.48)	7/1352
		<u>Streptomyces philanthi triangulum^T</u>	90.6/(99.45)	7/1265

Table S2. Complete 16S rRNA sequences of three Streptomyces spp.

>Streptomyces sp. B-81 Accesion Number NCBI (MW562807)

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Journal xxxxx 2022, 20, x FOR PEER REVIEW

CTGGCGGCGTGCTTAACACATGCAAGTCGAACGATGAACCACTTCGGTGGGGATTAGTGGCGAACGGGTGAG	510
TAACACGTGGGCAATCTGCCCTGCACTCTGGGACAAGCCCTGGAAACGGGGTCTAATACCGGATATTGACCTT	511
CACGGGCATCTGTGAGGTTCGAAAGCTCCGGCGGTGCAGGATGAGCCCGCGGCCTATCAGCTTGTTGGTGAGG	512
TAATGGCTCACCAAGGCGACGACGGGTAGCCGGCCTGAGAGGGCGACCGGCCACACTGGGACTGAGACACGG	513
CCCAGACTCCTACGGGAGGCAGCAGTGGGGGAATATTGCACAATGGGCGAAAGCCTGATGCAGCGACGCCGCG	514
TGAGGGATGACGGCCTTCGGGTTGTAAACCTCTTTCAGCAGGGAAGAAGCGAAAGTGACGGTACCTGCAGAA	515
GAAGCGCCGGCTAACTACGTGCCAGCAGCCGCGGTAATACGTAGGGCGCAAGCGTTGTCCGGAATTATTGGG	516
CGTAAAGAGCTCGTAGGCGGCTTGTCACGTCGGTTGTGAAAGCCCGGGGCTTAACCCCGGGTCTGCAGTCGAT	517
ACGGGCAGGCTAGAGTTCGGTAGGGGAGATCGGAATTCCTGGTGTAGCGGTGAAATGCGCAGATATCAGGAG	518
GAACACCGGTGGCGAAGGCGGATCTCTGGGCCGATACTGACGCTGAGGAGCGAAAGCGTGGGGGGGG	519
GGATTAGATACCCTGGTAGTCCACGCCGTAAACGGTGGGCACTAGGTGTGGGCAACATTCCACGTTGTCCGTG	520
CCGCAGCTAACGCATTAAGTGCCCCGCCTGGGGAGTACGGCCGCAAGGCTAAAACTCAAAGGAATTGACGGG	521
GGCCCGCACAAGCGGCGGAGCATGTGGCTTAATTCGACGCAACGCGAAGAACCTTACCAAGGCTTGACATAC	522
ACCGGAAACGGCCAGAGATGGTCGCCCCCTTGTGGTCGGTGTACAGGTGGTGCATGGCTGTCGTCAGCTCGTG	523
TCGTGAGATGTTGGGTTAAGTCCCGCAACGAGCGCAACCCTTGTCCCGTGTTGCCAGCAAGCCCCTTCGGGGG	524
TGTTGGGGACTCACGGGAGACCGCCGGGGTCAACTCGGAGGAAGGTGGGGACGACGTCAAGTCATCATGCCC	525
CTTATGTCTTGGGCTGCACACGTGCTACAATGGCCGGTACAATGAGCTGCGATACCGCAAGGTGGAGCGAATC	526
TCAAAAAGCCGGTCTCAGTTCGGATTGGGGTCTGCAACTCGACCCCATGAAGTCGGAGTCGCTAGTAATCGCA	527
GATCAGCATTGCTGCGGTGAATACGTTCCCGGGCCTTGTACA	528
>Streptomyces sp 796.1 Accession Number (MG654686)	529
GATTCCCTGGCTCAGGACGAACGCTGGCGGGCGTGCTTAACACATGCAGTCGAACGATGAGCCGCTTCGGTGG	530
TGGATTAGTGGCGAACGGGTGAGTAACACGTGGGCAATCTGCCCTGCACTCTGGGACAAGCCCTGGAAACGG	531
GGTCTAATACCGGATATGACTACCGATCGCATGGTTGGTGGTGGAAAGCTCCGGCGGTGCAGGATGAGCCCGC	532
GGCCTATCAGCTTGTTGGTGGGGTGATGGCCTACCAAGGCGACGACGGGTAGCCGGCCTGAGAGGGCGACCG	533
GCCACACTGGGACTGAGACACGGCCCAGACTCCTACGGGAGGCAGCAGTGGGGAATATTGCACAATGGGCGA	534
AAGCCTGATGCAGCGACGCCGCGTGAGGGATGACGGCCTTCGGGTTGTAAACCTCTTTCAGCAGGGAAGAAG	535

CGCAAGTGACGGTACCTGCAGAAGAAGCGCCGGCTAACTACGTGCCAGCAGCCGCGGTAATACGTAGGGCGC

GAGCGTTGTCCGGAATTATTGGGCGTAAAGAGCTCGTAGGCGGCTTGTCGCGTCGGATGTGAAAGCCCGGGGC

TTAACTCCGGGTCTGCATTCGATACGGGCAGGCTAGAGTTCGGTAGGGGAGATCGGAATTCCTGGTGTAGCGG

TGAAATGCGCAGATATCAGGAGGAACACCGGTGGCGAAGGCGGATCTCTGGGCCGATACTGACGCTGAGGAG

CGAAAGCGTGGGGAGCGAACAGGATTAGATACCCTGGTAGTCCACGCCGTAAACGTTGGGAACTAGGTGTGG

GCGACATTCCACGTCGTCCGTGCCGCAGCTAACGCATTAAGTTCCCCGCCTGGGGAGTACGGCCGCAAGGCTA

AAACTCAAAGGAATTGACGGGGGCCCGCACAAGCGGCGGGGGCATGTGGCTTAATTCGACGCAACGCGAAGAA

CCTTACCAAGGCTTGACATACACCGGAAACGGCCAGAGATGGTCGCCCCCTTGTGGTCGGTGTACAGGTGGTG

GCCAGCATGCCTTTCGGGGTGATGGGGACTCACAGGAGACTGCCGGGGTCAACTCGGAGGAAGGTGGGGACG

ACGTCAAGTCATCATGCCCCTTATGTCTTGGGCTGCACACGTGCTACAATGGCCGGTACAATGAGCTGCGATG

CCGTGAGGTGGAGCGAATCTCAAAAAGCCGGTCTCAGTTCGGATTGGGGTCTGCAACTCGACCCCATGAAGTC

GGAGTCGCTAGTAATCGCAGATCAGCATTGCTGCGGTGAATACGTTCCCGGGCCTTGTAACACA

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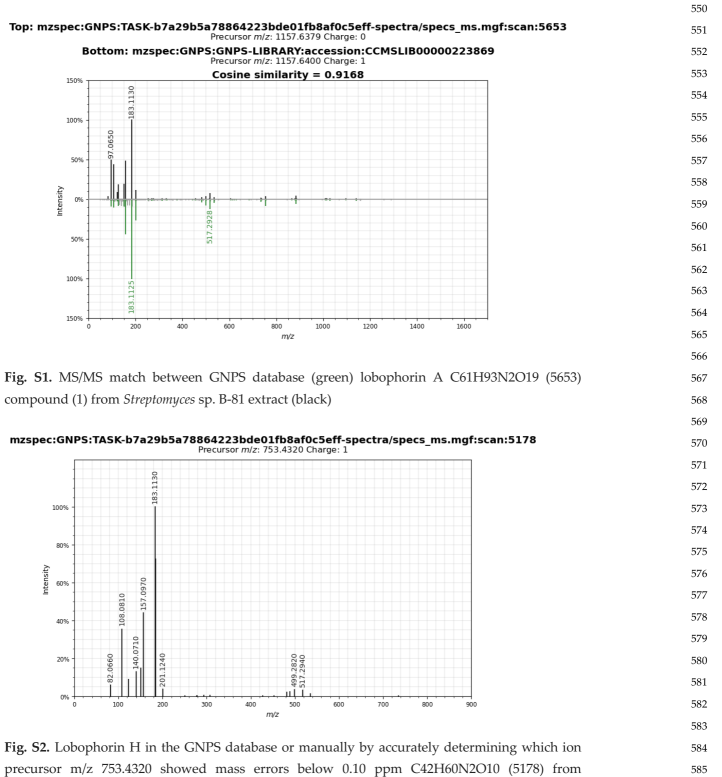
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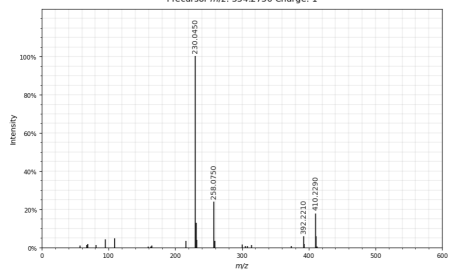
546

Journal xxxxx **2022**, 20, x FOR PEER REVIEW

Streptomyces sp. B-81 extract.

16 of 19





mzspec:GNPS:TASK-b7a29b5a78864223bde01fb8af0c5eff-spectra/specs_ms.mgf:scan:4097 Precursor m/z: 554.2750 Charge: 1

Fig. S3. Divergolide A in the GNPS database or manually by accurate determination of which ion precursor m/z 554.2750 showed mass errors below 0.10 ppm C31H39NO8 (4097) compound (3) from *Streptomyces* sp. B-81 extract.

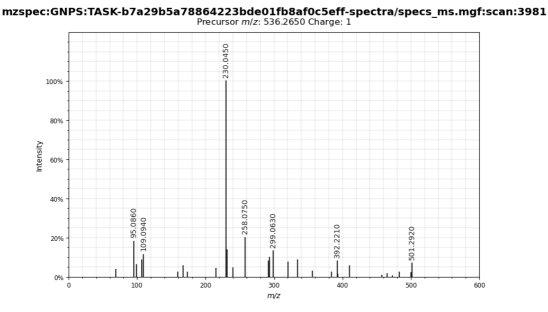
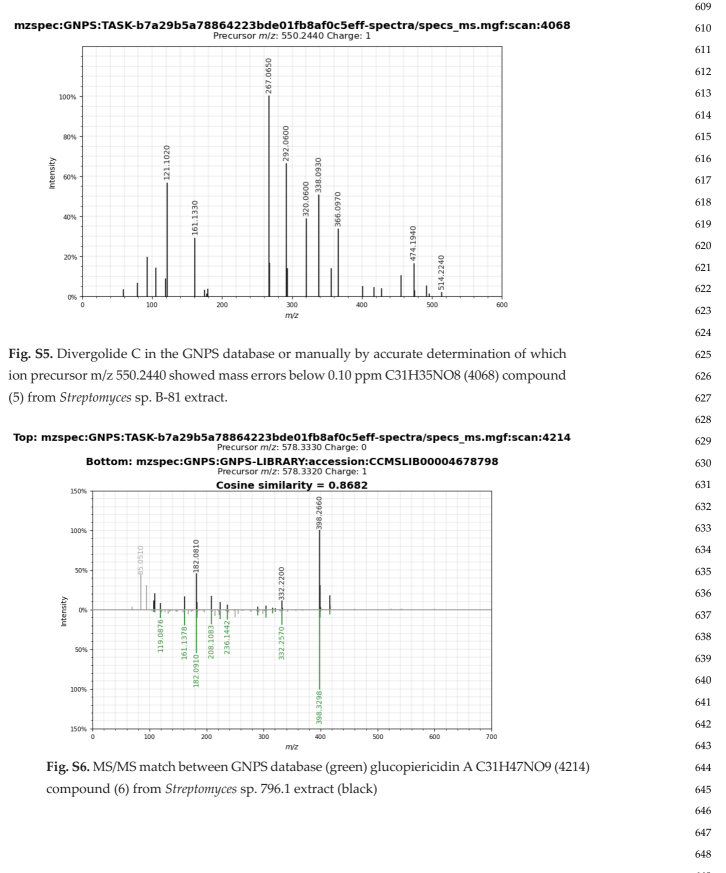


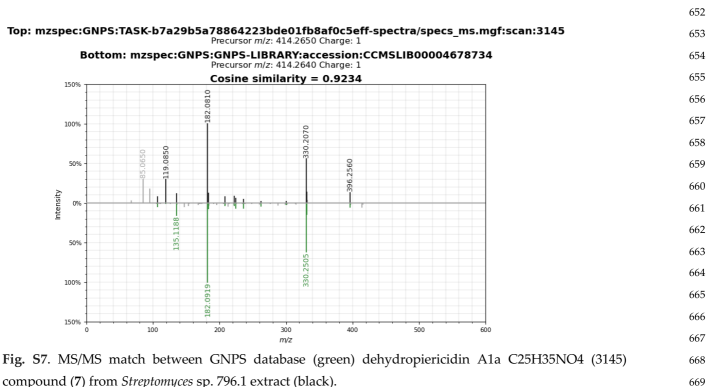
Fig. S4. Divergolide B in the GNPS database or manually by accurate determination of which ion precursor m/z 536.2650 showed mass errors below 0.10 ppm C31H37NO7 (3981) compound (4) from Streptomyces sp. B-81 extract.

Journal xxxxx 2022, 20, x FOR PEER REVIEW





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compound (7) from *Streptomyces* sp. 796.1 extract (black).