1	Discovery of a polybrominated aromatic
2	secondary metabolite from a planctomycete
3	points at an ambivalent interaction with its
4	macroalgae host
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32 Abstract

33 The roles of the majority of bacterial secondary metabolites, especially those from uncommon sources are yet elusive even though many of these compounds show striking biological activities. To further investigate the 34 secondary metabolite repertoire of underexploited bacterial families, we chose to analyze a novel representative 35 of the yet untapped bacterial phylum Planctomycetes for the production of secondary metabolites under 36 laboratory culture conditions. Development of a planctomycetal high density cultivation technique in 37 38 combination with high resolution mass spectrometric analysis revealed Planctomycetales strain 10988 to produce 39 the plant toxin 3,5 dibromo p-anisic acid. This molecule represents the first secondary metabolite reported from any planctomycete. Genome mining revealed the biosynthetic origin of this doubly brominated secondary 40 metabolite and a biosynthesis model for the com-pound was devised. Comparison of the biosynthetic route to 41 42 biosynthetic gene clusters responsible for formation of polybrominated small aromatic compounds reveals evidence for an evolutionary link, while the compound's herbicidal activity points towards an ambivalent role of 43 the metabolite in the planctomycetal ecosystem. 44

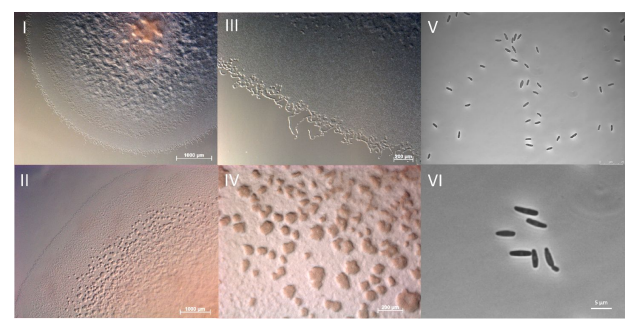
45 **1** Introduction

Bacterial secondary metabolism has long been a source of chemically diverse and biologically active natural 46 products.^{1,2} In fact, large numbers of biologically active entities have been isolated from extensively screened 47 phyla such as actinobacteria, firmicutes and proteobacteria.^{3–5} To establish alternative sources, natural products 48 49 research is increasingly focusing on taxa that have been less exploited to date, but show potential for production 50 of secondary metabolites according to the presence of secondary metabolite biosynthesis gene clusters (BGCs) in their genomes.² This strategic shift towards new producers increases chances for the discovery of novel bioactive 51 52 secondary metabolite scaffolds that are chemically distinct from the scaffolds found in previously screened bacteria. While it has long been stated that phylogenetically distant species have a more distinct secondary 53 54 metabolism, recent comprehensive secondary metabolome studies were able to validate this claim.^{6,7} Accordingly, it is now widely recognized that there is an urgent need to scrutinize novel bacterial taxa, alongside 55 the use of sensitive mass spectrometry and varied cultivation conditions to unearth novel natural products from 56 bacterial secondary metabolomes.⁸ Planctomycetes represent an underexploited phylum of bacteria in terms of 57 their secondary metabolite potential.⁹ Although planctomycetes have been already discovered in 1924, until now 58 59 no secondary metabolite of planctomycetal origin has been reported.¹⁰ This bleak picture is in clear contrast to 60 previous in-silico genome analysis that suggested planctomycetes to contain a significant number of secondary metabolite biosynthetic gene clusters (BGCs).^{9,11} In this work, we describe the first secondary metabolite from 61 any planctomycete including its structural characterization and biosynthesis, whereas its biological activity sheds 62 63 light on the putative ecological role within the planctomycete's natural habitat.

64 **2** Results and Discussion

2.1 Cultivation of *Planctomycetales* strain 10988

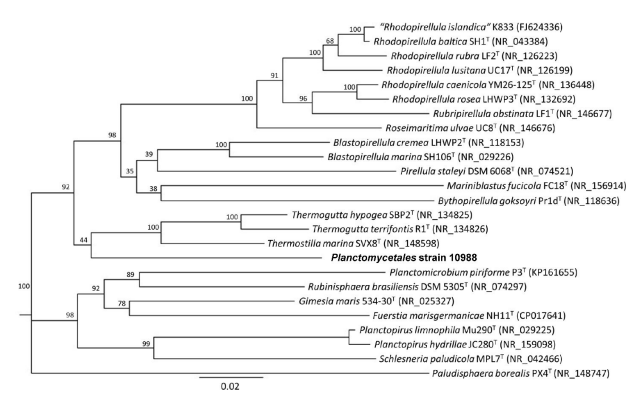
In order to investigate the biosynthetic capacity of uncommon and underexploited bacteria, we set out to
isolate new strains from marine sediment samples. Our efforts revealed a swarming, rose colored bacterial isolate
that was designated as strain 10988 (see Figure 1).



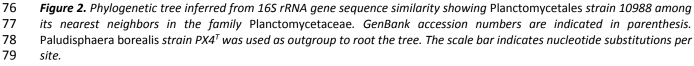
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Figure 1. Growth characteristics of Planctomycetales strain 10988 on solid medium displaying swarming (I and III), fruiting
 body like aggregate formation (II to IV) and phase contrast microscopy images of single cells from swarm (V) and from fruiting
 body like aggregates (VI).

- 72 16S rRNA-gene based phylogenetic analyses revealed the bacterium to belong to the order *Planctomycetales*,
- vhile being genetically distant from previously characterized planctomycetal genera such as Thermogutta,
- 74 Thermostilla, Pirellula, Rhodopirellula and Blastopirellula (see Figure 2).^{9,12–14}



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80 The nearest neighbors of strain 10988 among the Planctomycetales are the genera Thermogutta and Thermostilla, both of which are thermophilic as well as anaerobic or microaerophilic. However, strain 10988 81 82 showed very different characteristics. It does not only display optimum growth at 24 to 37 °C, but it also grows 83 aerobically. These findings agree with our 16S rDNA gene classification attempt, which classified strain 10988 as only distantly related to all characterized *Planctomycetes*. We thus sought to evaluate its secondary metabolome 84 85 as a potential source for new natural products. Although *Planctomycetes* have been continuously studied since 86 the late 1980s and Planctomycetes have shown to possess cellular features completely distinct from other prokaryotes, little has been done to investigate the planctomycetal secondary metabolome.^{15,16} One of the key 87 88 limiting factors that need to be overcome in order to uncover planctomycetal secondary metabolism is the 89 requirement to develop suitable cultivation techniques first. While there has been some success with cultivation 90 of freshwater *Planctomycetes*, cultivation of marine species such as strain 10988 turned out to be challenging.¹⁷ 91 Cultivation of strain 10988 showed that it is an obligate halophile, as it did not grow in absence of sea salts. The 92 halophilicity of the strain is underpinned by an ectoine biosynthesis gene cluster present in genome of strain 93 10988 that serves as a means to counterbalance the osmotic stress exerted by sea water brine on the cell. Furthermore, the strain depends on surface adsorption for efficient growth which is exemplified by its enhanced 94 growth in early stages if cellulose powders are added to the medium. As these fine filter paper pieces turn to rose 95 96 color before the medium in the shake flask contains a significant number of suspended cells, the bacterium seems to preferentially colonize surfaces before dispersing into suspension in a shake flask (supporting information). 97

98 This finding is well in line with planctomycetal growth in nature that occurs in parts fixed to the surfaces of macroalgae.⁹ The slow growth of this isolate in combination with initially low secondary metabolite production 99 100 rates – as judged by LC-MS analysis - led us to devise a fermenter based cultivation to obtain increased secondary 101 metabolite yields that could not be achieved in shake flask cultivations. As a means to stimulate productivity of 102 the planctomycetal strain for secondary metabolite isolation, we added adsorber resin to shake flask cultures. 103 This should circumvent productivity limitations arising from feedback inhibition mechanisms.¹⁸ However, addition 104 of adsorber resin led to complete suppression of planctomycetal growth unless the culture was inoculated with a 105 high concentration of actively growing cells. When strain 10988 was grown in absence of adsorber resin, 106 inoculation of liquid cultures with a very low concentration of live cells was sufficient to stimulate planctomycetal 107 growth. The most probable explanation for this phenomenon is that the presence of adsorber resin in low density 108 cultures masks certain quorum sensing signals by binding them, inhibiting cooperative growth of planctomycetes. 109 While quorum sensing has been linked to different effects such as the inhibition of biofilm formation or virulence, 110 a guorum sensing signal that increases or stalls cell division speed has not been described yet.¹⁹ As a result, in 111 order to avoid lack of growth or unnecessarily lengthy lag phases in planctomycetal fermentations in larger scale 112 production cultures, adsorber resin addition was performed several days post inoculation of the respective 113 fermenter. In analytical scale shake flask cultivations, the effect of adding resin directly could be mitigated by 114 inoculation of the cultures with a higher concentration of live cells.

2.2 Discovery of 3,5-dibromo p-anisic acid

115 To assess the secondary metabolome of planctomycetal strain 10988, methanolic extracts of the strain's 116 culture supplemented with adsorber resin were prepared and compared to methanolic extracts of the corresponding medium ("blank" sample) to obtain an overview about its secondary metabolome. The bacterial 117 118 extract as well as the blank were subjected to high-resolution LC-MS analysis using a reverse-phase UPLC-119 coupled qTOF setup (supporting information). This analysis revealed an intriguing signal presenting a monoisotopic mass of 308.873 Da $[M+H]^+$ (C₈H₆Br₂O₃, Δ m/z = 6 ppm) and an isotope pattern that pointed 120 towards double bromination.²⁰ Maximum cell density as well as the production rate of this doubly brominated 121 122 compound remained limiting for material supply in shake flask cultures even after media optimization 123 (supporting information). We therefore developed a method to grow strain 10988 in a fermenter which allowed 124 to purify the candidate compound by semi-preparative HPLC (supporting information).

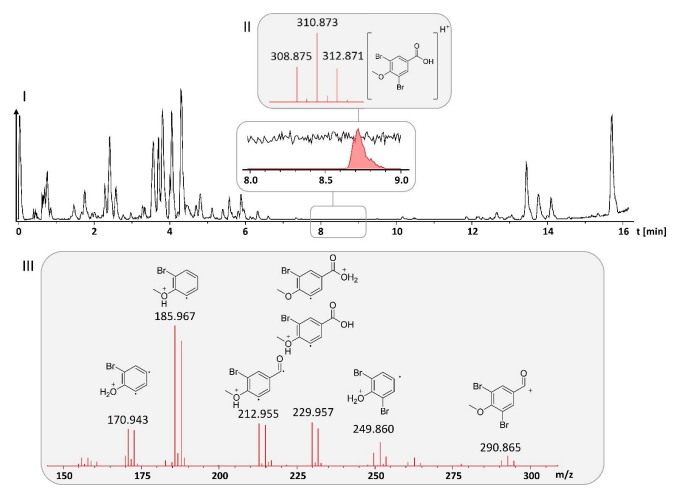


Figure 3. I) LC-MS chromatogram of Planctomycetales strain 10988 with magnification of the MS signal for 3,5 dibromo p anisic acid (1) II) Corresponding MS spectrum and structure formula of 1 III) MS² spectrum of 1 and putative product ions
 formed in MS² fragmentation.

According to molecular formula calculation the molecule possesses 6 hydrogen atoms. As the ¹H NMR 129 spectrum contains only 2 singlet signals, the molecule has to be highly symmetrical. One of the signals 130 131 representing 2 protons has a strong downfield shift of 8.11 ppm indicating a heavily electron deficient symmetrical aromatic system while the other singlet signal representing 3 protons is characteristic for an oxygen-linked methyl 132 group. The ¹³C shift of the corresponding methyl group indicates its connection to the phenolic oxygen of the 133 134 molecule and not to the carboxylic acid (supporting information). This is further supported by the fact that the 135 tandem MS spectra show a strong water loss as expected from free carboxylic acid moieties, while we did not observe neutral loss of methanol in tandem MS experiments (Figure 3). This neutral loss would be expected if the 136 molecule contained a methyl ester. The double brominated compound produced by 10988 was therefore 137 138 determined to be 3,5 dibromo-p-anisic acid (1), which could be later confirmed using synthetic standard material. 139

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2.3 Biosynthesis of 3,5-dibromo p-anisic acid

141 In order to identify the biosynthetic origin of 1, we determined the complete genome of *Planctomycetales* 142 strain 10988 using PacBio long read sequencing technology (supporting information). Genome assembly resulted 143 in a single circular bacterial chromosome of 6.6 Mbp with a total GC content of 50.4% (GenBank accession number 144 XXX). AntiSMASH analysis of the bacterial genome annotated 3 terpene biosynthetic gene clusters (BGCs), an 145 ectoine BGC, a cluster for lassopeptide biosynthesis and a PKS type III BGC.²¹ Contrary to earlier genome mining 146 results from planctomycetes, the genome of strain 10988 does not encode any multimodular secondary metabolite pathways in its genome.^{11,22} Since our newly elucidated secondary metabolite is likely not produced 147 by a multimodular megasynthase, the biosynthesis gene cluster predictors run by antiSMASH are in this case 148 149 unsuitable to annotate the corresponding biosynthesis pathway, which possibly consists of a set of 'stand-alone' 150 enzymes.²¹ We therefore searched the obtained genome for flavin dependent halogenase enzymes, as most 151 region selective bromination or chlorination reactions on aromatic systems are catalyzed by this protein family in 152 nature. ^{23,24} The flavoprotein showing highest homology to halogenating enzymes was named BaaB. It was found 153 encoded adjacent to - and putatively on the same mRNA strand as - a chorismate lyase-like protein termed BaaA. This protein could plausibly deliver the precursor para-hydroxy benzoeic acid from the cellular chorismic acid pool 154 (Figure 4).²⁵ Homology modelling of the two proteins on the protein fold recognition server Phyre2 supports this 155 finding, as both proteins involved in biosynthesis of **1** are correctly mapped onto the expected protein families.²⁶ 156 157 Unfortunately, and despite serious efforts we were unable to develop methods to genetically manipulate the 158 planctomycetal strain and we were thus unable to further validate our hypothesis by an inactivation mutant of 159 the baaA and baaB locus. The mechanism of UbiC-like chorismate lyases such as BaaA is less studied in comparison 160 to the mechanism of chorismate mutases. Chorismate mutase reactions consist of an electrocyclic 6 electron rearrangement reaction that leads to prephenate formation (Figure 4).²⁵ Chorismate lyase enzymes like BaaA use 161 162 a closely related electrocyclic 6 electron rearrangement reaction that removes pyruvate from chorismate to form 163 p-hydroxy benzoeic acid.²⁷ At this point we cannot differentiate whether p-hydroxybenzoeic acid is methylated 164 to p-anisic acid first or if methyl transfer occurs after double bromination of p-hydroxybezoeic acid by the 165 brominase enzyme BaaB. In order to finish biosynthesis of 1 after the action of BaaA and BaaB, an SAM dependent 166 O-methyl transferase (BaaC) is needed that transforms 2 into 1. When analyzing the genetic locus encoding BaaA 167 and BaaB we did not find such an enzyme, meaning BaaC is encoded in a different genetic locus. The fact that the 168 baaA and baaB genes are encoded adjacently in the genome, while the corresponding methyl transferase baaC 169 is encoded in a different location may indicate that dibromo p-hydroxybenzoeic acid is produced fist and 170 subsequently methylated. It is worth noting that we could not identify either of the possible intermediates via LC-171 MS from the 10988 strain extracts. As BaaB is the only halogenase enzyme encoded in the baa BGC, it is certainly 172 responsible for 3,5 dibromination of the aromatic moiety. As both positions that are brominated are chemically 173 equivalent it is not surprising that both halogenations are performed by the same enzyme. Furthermore, we 174 observe strict specificity of BaaB for bromine as no chlorinated or mixed brominated and chlorinated anisic acid

derivatives can be identified in the fermentation broth. Thus, BaaB is either unable to bind chloride anions instead 175 176 of bromide anions due to a difference in binding cavity size, or the redox potential of BaaB is not sufficient to 177 oxidize chloride anions but is sufficient to oxidize bromide ions to an activated species. Still, BaaB is not the only 178 such enzyme unable to process chlorine, as the brominase Bmp5 from Pseudoalteromonas strains involved in biosynthesis of polybrominated phenols is also specific for bromine over chlorine.²⁸ The architectures of the 179 responsible loci producing polybrominated biphenylic secondary metabolites show remarkable similarity to the 180 181 Baa operon even though the host organisms are phylogenetically very distant. While Bmp5-like proteins from P. luteoviolacea 2ta16 and P. phenolica O-BC30 are very similar as they share 96% homology, their similarity to BaaB 182 183 remains around 44%. This finding is readily explained as Pseudoalteromonads and Planctomycetes are 184 phylogenetically far apart and BaaB only catalyzes meta-position bromination, while Bmp5 also catalyzes ipso 185 substitution of CO₂ at the aromatic core. This reaction removes the carboxylic acid and the phenols are not 186 afterwards.^{29,30} Furthermore, in biosynthesis of polybrominated biphenyl ethers in methylated 187 Pseudoalteromonads, an additional enzyme called Bmp7 uses phenolic coupling reactions to form biphenyl 188 structures that do not exist in our planctomyces strain, as the baa gene cluster in strain 10988 does not possess the corresponding CYP P450 enzymes. ³⁰ The absence of CYP enzymes in the planctomycetal BGC explains why 189 190 the planctomycete only synthesizes monocyclic polybrominated aromatic compounds, as it lacks the CYP enzyme required to perform phenol couplings leading to the formation of biphenylic compounds.^{30,31} 191

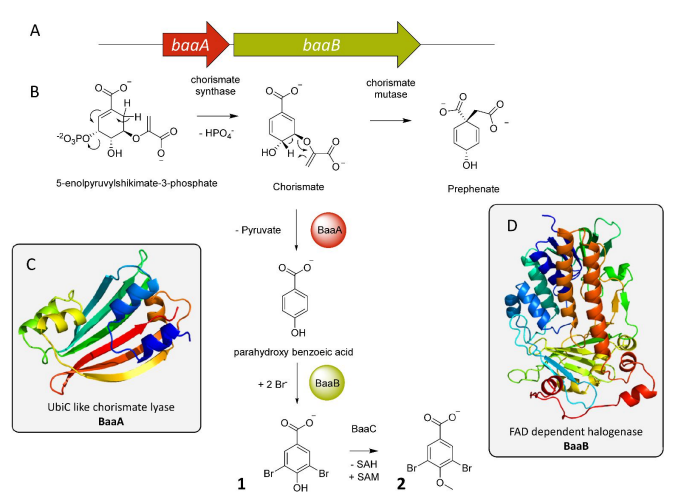


Figure 4. A) Gene cluster for 3,5 dibromo p-anisic acid (1) biosynthesis B) Proposed model for biosynthesis of 1 including the
 non-methylated precursor 2 C) Phyre2 based homology model for the chorismate lyase BaaA D) Phyre2 based homology model
 for the brominase BaaB.

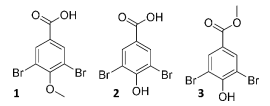
197 As shown in Figure 4, in order to finish biosynthesis of 1, the formerly mentioned oxygen-methyl transferase 198 BaaC is required that transfers a methyl group and thus forms the methoxy group in **1**. The corresponding oxygen-199 methyl transferase responsible for methyl transfer to the phenolic oxygen of the precursor compound is not 200 clustered with the genes responsible for p-anisic acid production. Although there are some candidate sequences 201 for the BaaC protein, it is impossible to exactly pinpoint the methyl transferase in the 10988 genome responsible for methylation of 2 due to our inability to perform directed mutagenesis with the strain. Nevertheless, following 202 203 blast analysis of the 10988 genome for homologues to the StiK protein from stigmatellin biosynthesis in S. aurantiaca (NCBI protein acc. Nr. CAD19094.1) and UbiG from Escherichia coli K12 (NCBI protein acc. Nr. 204 205 BAA16049.1) that both catalyze methyl transfer to an aromatic hydroxyl group, we obtained 8 candidate 206 sequences. ^{32,33} These candidate genes for the enzyme BaaC, which show similarity to both aforementioned enzymes, can thus be assumed to catalyze reactions such as the transformation of 2 to 1. 207

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2.4 Bioactivity evaluation of 1 and its analogs

Due to the double bromination of **1**, its biosynthesis is costly to the strain in terms of energy and resources. Therefore, **1** can be plausibly expected to confer a competitive advantage to strain 10988 in its environment. To evaluate this biological role of **1** we set out to profile its bioactivity as well as the bioactivity of its biological precursor **2** and its isomer Methyl 3,5 p-hydroxy benzoeic acid (**3**). The compounds **2** and **3** that cannot be obtained from the planctomycetal culture broth are commercially available.



Scheme 1. The natural product 3,5 dibromo p-anisic acid (1), its putative biological precursor 3,5 dibromo p-hydroxybenzoeic
 acid and the natural product analog Methyl-3,5 dibromo p-hydroxybenzoate (3)

The planctomycetal natural product 1, its precursor 2 and its analog 3 were tested in an antibiotics screening 218 against the bacterial pathogens C. freundii, A. baumanii, E. coli, M. smegmatis, S. aureus, P. aeruginosa, B. subtilis 219 220 and M. luteus, the yeasts C. albicans and P. anomala as well as the filamentous fungus M. hiemalis. The compounds did not display any inhibition of these microbial indicator strains at concentrations up to 64 μ g/ml. 221 To evaluate cytotoxicity we tested 1, 2 and 3 in a cell line cytotoxicity assay. This assay revealed both methylated 222 compounds (1 and 3) to display moderate cytotoxicity to the human cervical carcinoma cell line KB3.1. While 1 223 224 and **3** showed an IC₅₀ value of 60 μ g/ml, no cytotoxicity was found for the free acid **2** μ p to 300 μ g/ml. To assay herbicidal activity of p-methoxy dibromo benzoeic acid as well as its precursor and isomer we tested the 225 226 compound's activity on the germination of Agrostis stolonifera penncross. IC₅₀ values for seed germination 227 inhibition were determined to be 32 μ g/ml for **1**, 64 μ g/ml for **2** and 16 μ g/ml for **3**. The fully decorated methoxy 228 derivative 1 shows higher biological activity than its precursor 2 while the non-natural methyl ester derivative 3 229 shows the best anti-germination activity in the A. stolonifera germination assay (supporting information). As 230 compound **1** shows no significant antibacterial activity or mammalian cell cytotoxicity but displays moderate 231 phytotoxicity and as 1 is likely released to the marine environment, we assume its role as a putative algal toxin 232 (supporting information).

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2.5 On the potential biological role of 1

238 Many planctomycetes live in close association with macroalgae which they cover almost completely.³⁴ Given 239 that the strain 10988 was isolated from a marine sediment, the strain is likely associated with marine algae in its 240 natural habitat. The exact mode of interaction between the planctomycetes and the macroalgae has yet to be 241 determined. Still, the high abundance of planctomycetes on algal species, which can reach up to 50 % of the algal 242 microbiome, indicates that these bacteria are significant interaction partners for the algae.³⁴ One hypothesis 243 considers the algae as a food source for planctomycetes, since they are able to utilize uncommon sugars such as 244 rhamnose and fucose contained in algal biomass. In our case, strain 10988 – like other planctomycetes – was able 245 to grow on uncommon sugars such as galactose, mannose, lactose, sucrose, maltose raffinose, xylose and rhamnose (supporting information, Figure S2), indicating that this nutritional option could apply to strain 10988. 246 ¹¹ As on the other hand, the planctomyces bacterium possesses the ability to produce the plant toxin 3,5 dibromo 247 248 p-anisic acid, whose production seems to be tightly controlled as judged by the low production titers under 249 laboratory conditions, we reason that an ambivalent interaction model might take place between planctomycetes 250 and their plant hosts. The planctomycetes strains probably live on the algal surface and modulate the local microbial community until they sense the algal species they live on is weakening. This might trigger expression of 251 said plant toxin to kill and decay this part of the algae and the bacterium would subsequently move on to colonize 252 253 different algae. Similar 'Jeckyll and Hyde' behavior, meaning the ability to switch between commensalism and 254 symbiosis, comparable to planctomycetal colonization of algae, and a virulent state that is hostile to its host 255 organism has been described for the human pathogen C. albicans.³⁵ The ability of strain 10988 to produce a plant 256 toxin as a bacterium associated to macroscopic plants can be seen as a strong hint that the bacterium adopts such 257 a strategy.

258 **3** Conclusion

259 In this work we describe the cultivation of a new marine planctomycete that is genetically distant from all 260 planctomycetes known to date, and reveal Planctomycetales strain 10988 as producer of a dibrominated 261 secondary metabolite. Isolation of this secondary metabolite required a stirred tank reactor setting and optimized 262 medium and culture conditions. Subsequent structure elucidation of 1 by NMR revealed an intriguing structure 263 and thus sparked interest in the biosynthetic origin and ecological role of the compound. We were able to pinpoint 264 the core biosynthesis genes baaA and baaB that can accomplish the core structure of 1. Investigation of the 265 bioactivity of **1** as well as the bioactivity of its isomer **3** and putative precursor **2** showed that this compound class 266 displays herbicidal activity in A. stolonifera penncross germination assays, leading us to hypothesize on a biological 267 role of this compound in the life cycle of the algal symbiont *Planctomycetales* strain 10988. In conclusion, we contribute to the understanding of the biogenesis of small polyhalogenated compounds in marine bacteria, 268

whereas it remains astonishing to what extent such polybrominated aromatic substances are apparently released
 into the ecosystem from biological instead of anthropogenic sources.

271 This study also identifies planctomycetes as an underexploited source of biologically active secondary metabolites, as **1** to the best of our knowledge is the first natural product described from this bacterial taxon. 272 However, we would like to point out that previous studies may have overestimated the genome encoded 273 secondary metabolite diversity of planctomycetes as a group, since strain 10988 under study here did not show 274 the presence of any multimodular megasynthetase.^{11,22} Even though strain 10988 shows some BGCs, especially 275 BGCs linked to terpene biosynthesis, megasynthase containing BGCs are often considered as benchmark indicators 276 277 for secondary metabolite production capability.² On the other hand, the example presented here shows how important it is to evaluate new taxa on the metabolomics stage, since metabolites of the type described here are 278 279 easily missed by genome mining. Thus, the overall potential of planctomycetes awaits further investigation. While 280 growing these bacteria under laboratory conditions may be tedious and non-trivial, devising methods for their 281 cultivation is a valuable tool to tap into the planctomycetal secondary metabolite space. The discovery of 282 polybrominated compounds in strain 10988 is well in line with both the observation that this bacterium is an 283 obligate halophile and reminiscent of the strain's marine origin. The isolated and characterized natural product 284 3,5 dibromo p-anisic acid shows that Nature, especially the marine microbial community is able to biosynthesize 285 polyhalogenated small aromatic compounds that look like anthropogenic products of chemical synthesis.

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292 **Supporting Information**

An in detail description of the planctomyces strain, all utilized fermentation protocols, *in silico* analyzes on gene and protein level as well as all relevant NMR data for structure elucidation are available free of charge via the Internet at BioRxiv.org.

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