Sponge-Inspired Dibromohemibastadin Prevents and Disrupts Bacterial Biofilms without Toxicity

Tiffany Le Norcy 1, Hendrik Niemann 2, Peter Proksch 2, Karen Tait 3, Isabelle Linossier 1, Karine Réhel 1, Claire Hellio 4,* and Fabienne Fay 1

1 Laboratoire de Biotechnologie et Chimie Marines, Institut Universitaire Européen de la Mer, Université de Bretagne-Sud, 56100 Lorient, France; tiffany.le-norcy@univ-ubs.fr (T.L.N.); isabelle.linossier@univ-ubs.fr (I.L.); karine.rehel@univ-ubs.fr (K.R.); fabienne.fay@univ-ubs.fr (F.F.)
2 Institute of Pharmaceutical Biology and Biotechnology, Heinrich-Heine-University Düsseldorf, 40225 Düsseldorf, Germany; hendrik-niemann@web.de (H.N.); peter.proksch@hhu.de (P.P.)
3 Plymouth Marine Laboratory, Plymouth PL1 3DH, UK; KTAIT@pml.ac.uk
4 Biodimar, LEMAR UMR 6539, Institut Européen de la Mer, Université de Bretagne Occidentale, 29200 Brest, France
* Correspondence: Claire.hellio@univ-brest.fr

Received: 2 June 2017; Accepted: 6 July 2017; Published: 12 July 2017

Abstract: Since the banning of several families of compounds in antifouling (AF) coatings, the search for environmentally friendly AF compounds has intensified. Natural sources of AF compounds have been identified in marine organisms and can be used to create analogues in laboratory. In a previous study, we identified that dibromohemibastadin-1 (DBHB) is a promising AF molecule, leading to the inhibition of the activity of phenoloxidase, an enzyme involved in the attachment of mussels to surfaces. This paper describes the activity of the DBHB on biofilm formation and its detachment and on bacterial adhesion and communication: quorum sensing. DBHB has an anti-biofilm activity without affecting adhesion of marine and terrestrial bacteria at a dose of 10 µM. Moreover, DBHB activity on quorum sensing (QS) is demonstrated at doses of 8 and 16 µM. The activity of DBHB on QS is compared to kojic acid, a quorum sensing inhibitor already described. This compound is a promising environmentally friendly molecule potentially useful for the inhibition of microfouling.

Keywords: biofouling; biofilm; bastadin derivative; microfouling; quorum sensing; sponge

1. Introduction

All surfaces immersed in aquatic environments are rapidly colonized by organisms (microfoulers and macrofoulers); this natural phenomenon is defined as biofouling [1].

It is occurring in three main sequential stages: immediately after immersion, surfaces are covered by dissolved organic molecules which then stimulate subsequent adhesion of microorganisms such as bacteria, fungi and microalgae; these organisms constitute the microfouling. It acts as attractant for the adhesion of macrofoulers such as macroalgae (zygote and/or spores) and invertebrate larvae [2–4].

Within biofilms, bacteria are usually present in higher densities than in other microorganisms. These bacterial communities are constituted by sessile bacteria, which are adhered to each other and/or to the surface, with the adhesion being facilitated by the production of exopolysaccharides (EPS) [5]. The formation of microbial biofilm is a complex process which involves communication such as quorum sensing (QS). QS is a microbial signaling system which allows for the regulation of population behaviors such as the production of virulence factors [6] or the formation of biofilm [7]. The QS system involves small molecules named autoinducers (AI). These molecules, linked to the protein receptor, induce the expression of several genes involved in population behaviors such as biofilm formation, motility or pigment production. One of the family of QS signal molecules are
acyl-homoserine lactone (AHL) type molecules. This bacterial communication was first described in a marine bacteria *Vibrio fisheri* [8]. Later, similar signaling systems were discovered in other bacterial species. Two systems are described: in Gram negative bacteria, AHL-like communication molecules are present, whereas in Gram positive bacteria, other molecules (autoinducer-2: pheromones) are identified [9]. It was described that QS plays an important role in numerous processes governing biofilm formation and organization. Waters et al., demonstrated that when *Vibrio cholera* was in low cell density, the phosphorylation of LuxO inhibited an mRNA which induced the expression of a protein inhibiting the C-di-GMP. The C-di-GMP, an intracellular messenger, has already been shown to be involved in bacterial biofilm formation [7]. In the same way, *P. aeruginosa* mutated on the lasI gene induced a biofilm 20% finer than the wild type [10]. The discovery of a molecule which could interfere with the QS and especially with the biofilm formation would be a promising candidate for new anti fouling (AF) compound [11] in replacement of current AF compounds used in paint, such as heavy metals or biocides.

Some compounds have attracted wide-spread attention for their ability to interact with QS. For example, molecules of the furanone family are able to interfere with bacterial communication [12,13]. These compounds possess a chemical structure similar to the structure of AHL [14]. Other molecules, some enzymes (lactonase or acylase) could be used to inhibit QS [15]. For example, an enzyme named AiiA is able to inhibit the QS by the degradation of the lactone ring of AHL, which is involved in the recognition of the receptor [16].

The environment is a source of inspiration for many innovations. In nature, numerous marine organisms which are fixed and/or moving slowly are able to inhibit biofouling [17,18]. Numerous bioactive substances have been isolated from marine organisms such as algae and sponges [19,20]. Such compounds contribute, in most cases, to the chemical defense against predation, or to the inhibition of adhesion and growth of pathogens and epibionts. Sponges are pre- eminent producers of bioactive secondary metabolites with antibacterial, antifungal or anti-inflammatory activities [21–23]. From natural molecules, it is also possible to synthesize analogues, which possess equivalent or even stronger activities, due to the addition of specific chemical groups [24–26].

Within this study, we decided to focus on the potential of bastadins as AF inhibitors. Bastadins have been previously isolated from a marine sponge *Ianthella basta*. Thirty natural bastadins were characterized and screened for their AF potencies: it was highlighted that some of these compounds were able to inhibit barnacle adhesion and adhesives synthesis by mussels [27,28]. Bastadins are composed of a brominated tyrosine and a brominated tyramine unit linked by a peptide bond in order to form a hemibastadin unit (Figure 1A). The tyrosine moiety furthermore bears an oxime function. Two hemibastadins linked by covalent bonds give rise to a bastadin. Within this study, we focused on the characterization of the activity of a hemibastadin analogue named dibromohemibastadin-I (DBHB) (Figure 1B). DBHB consists of a hemibastadin unit composed by brominated tyrosine and a likewise brominated tyramine unit; its synthesis was previously described by Bayer et al. in 2011 [27]. DBHB is selected and evaluated in our study because this compound has already been described as an inhibitor of macroorganisms (barnacles and mussels) adhesion [28,29] without toxicity. However, its activity on microorganisms has never been described. Microorganisms are important in biocorrosion, and these organisms can develop resistance against biocides, so it is important to identify new molecules and strategies to limit biofouling. Ortlepp et al. in 2007 published data about DBHB activity toward cyprid larvae of *Barnacle improvisus* and nauplii of *Artemia salina*: DBHB induced an inhibition of the larval settlement at 10 µM without mortality and did not cause mortality of the brine shrimp nauplii when used at 10 µM [29]. The importance of the oxime function and bromine atoms for AF activity was also highlighted. Bayer et al. in 2011, showed that DBHB inhibits phenoloxydase (enzyme involved in adhesive production by *Mytilus sp.*) activity with an IC$_{50}$ of 0.84 µM [27].
The aim of our work is to further characterize the activity of DBHB, with a focus on marine and terrestrial bacterial adhesion and biofilm formation. These experiments have been realized in dynamic conditions, in a flowcell system, established by Tolker-Nielsen [30]. The molecule has also been tested on bacterial communication to determine its mode of action.

2. Results

2.1. Anti-Bacterial Activity

To determine the antibacterial property of DBHB, the activity of the molecule was evaluated on the growth of four bacteria, three marine bacteria (Paracoccus sp. 4M6, Pseudoalteromonas sp. 5M6 and Vibrio sp. D66) isolated in the Gulf of Morbihan (south Brittany) [31] and a terrestrial bacterium Pseudomonas aeruginosa PAO1 used as reference. DCOIT was used as positive control, it is the active compound of the seanine® [32].

Results do not show any statistical differences (ANOVA, p > 0.05) unlike to the addition of the DCOIT, which induced an inhibition of the bacterial percentage by a factor of 4 (Figure 2). The inhibition of bacterial growth by the DCOIT is statistically different to the control (ANOVA, p < 0.01).

The anti-bacterial test showed that DBHB does not affect the bacterial growth at doses between 0.02 and 80 µM. DBHB is not toxic for the bacteria studied.

To evaluate the activity of the compound DBHB on bacterial adhesion and biofilm, the concentration tested varied from 2 to 16 µM. This range of concentration was selected for the next experiments because these concentrations were high enough for activity evaluation.
2.2. Impact of DBHB on AHL Production

Amongst the four strains studied (PAO1, 4M6, 4J6 and 5M6), three were shown to be able to produce AHLs. However, no AHL was identified in the supernatant of *Pseudoalteromonas* sp. 5M6. The AHL identification is presented in Table 1.

**Table 1.** Identification of acyl-homoserine lactones (AHLs) produced by *P. aeruginosa* PAO1, *Paracoccus* sp. 4M6, *Bacillus* sp. 4J6 and *Pseudoalteromonas* sp. 5M6.

| Bacterial Strains               | Gram  | AHL Produced                                      |
|--------------------------------|-------|--------------------------------------------------|
| *Pseudomonas aeruginosa* PAO1  | Negative | C₄-HSL, C₆-HSL and 3-oxo-C₁₂-HSL                   |
| *Paracoccus* sp. 4M6            | Negative | C₄-HSL, C₆-HSL, C₈-HSL and 3-oxo-C₁₀-HSL          |
| *Bacillus* sp. 4J6              | Positive | Autoinducer-2                                    |
| *Pseudoalteromonas* sp. 5M6     | Negative | No. AHL                                          |

The two bacteria gram negative *P. aeruginosa* PAO1 and *Paracoccus* sp. 4M6 produce three or four different AHLs. For the other gram negative bacterium, *Pseudoalteromonas* sp. 5M6, no AHL was identified. It is probable that this bacterium produces another kind of communication molecule. For the last bacterium, *Bacillus* sp. 4J6, in gram positive bacterium, other molecules of communication have been identified in the literature (Autoinducer-2, AI-2) [33,34].

The addition of DBHB does not modify the AHL production, chromatograms of *P. aeruginosa* PAO1, *Paracoccus* sp. 4M6; and *Pseudoalteromonas* sp. 5M6, with or without DBHB, were similar.

2.3. Anti-Adhesion Activity

DBHB activity was evaluated on the adhesion of the marine bacterium *Paracoccus* sp. 4M6 and of the terrestrial bacterium *Pseudomonas aeruginosa* PAO1. The molecule activity was determined in the flowcell system [30]. The addition of DBHB in both conditions (within the bacterial growth media or by conditioning the adhesion surface) did not induce any inhibition or activation of the adhesion of *P. aeruginosa* PAO1 or *Paracoccus* sp. 4M6 in comparison to the control. Figure 3 summarizes the observations obtained by Confocal laser scanning microscopy (CLSM) on the adhesion of *P. aeruginosa* PAO1 and *Paracoccus* sp. 4M6 on the glass slide.

**Figure 3.** Confocal laser scanning microscopy observations of the bacterial adhesion with syto9® after 2 h with addition of DBHB at 16 µM in two conditions (A: PAO1 control, D: 4M6 control, B,E: addition of DBHB to the bacterial suspension for PAO1 and 4M6, C,F: conditioning of the adhesion surface for PAO1 and 4M6); the overlap percentage (%) is added on observations, the ± represents the standard deviation.
It was also observed that the strain *P. aeruginosa* PAO1 adhered far more than the bacterium *Paracoccus* sp. 4M6, there was a factor of 4 between the overlap percentages of *P. aeruginosa* PAO1 and *Paracoccus* sp. 4M6. This difference of overlap percentage was statistically different ($p < 0.01$) and could be explained by the different ability of the bacteria to adhere.

### 2.4. Anti-Biofilm Activity

#### 2.4.1. Biofilm Formation

The impact of DBHB was observed on the biofilm formation. The activity of the molecule was tested on Gram negative bacteria, which produce AHL (*Paracoccus* sp. 4M6 and *P. aeruginosa* PAO1), on a gram negative bacteria and which did not produce AHL (*Pseudoalteromonas* sp. 5M6), as well as on a Gram positive bacterium (*Bacillus* sp. 4J6). AHL production was determined after an extraction and a LC–MS–MS dosage (Data are not shown) [35].

For the strain *P. aeruginosa* PAO1, observations obtained in CLSM are shown in Figure 4.

![Figure 4](image)

*Figure 4.* Observation of *P. aeruginosa* PAO1 biofilms (syto9®) with addition of DBHB at four concentrations (2, 8, 16 and 32 µM) in the medium growth flow, biovolumes (µm³/µm²) and biofilms percentages (%) are presented; the ± represents the standard deviation.

For *P. aeruginosa* PAO1, the inhibition of the biofilm was statistically different from the control only at 16 and 32 µM. At 2 and 8 µM the biofilm was inhibited by only a factor of 1.3. The standard deviations at 2 and 8 µM DBHB was higher than those observed for 16 and 32 µM DBHB, this explains why the biofilm inhibition was not statistically different after analysis with the ANOVA and Tukey tests. During these experiments, the strain *P. aeruginosa* PAO1 at times formed filaments in biofilm and these filaments could modify the biofilm structure and the biovolume associated. The modification of the biofilm structure in *P. aeruginosa* PAO1 could be linked to environmental factors (temperature, pH...). This is why the activity of biofilm inhibition is not the same for the different concentrations evaluated; in flowcell system, it was not possible to control the filament formation. The activity of the compound DBHB was evaluated towards a second strain, a Gram negative bacteria producing AHL: the marine bacterium *Paracoccus* sp. 4M6 (Figure 5).

For the strain *Paracoccus* sp. 4M6, the addition of DBHB to the growth medium induced an inhibition of the biofilm which was significantly different from the control at the four tested concentrations. The biovolumes of *Paracoccus* sp. 4M6 biofilm were 2.1, 2.6, 2.5 and 3.2 times lower than the control for the addition of DBHB at 2, 8, 16 and 32 µM respectively. The similarity of these two bacteria is their ability to produce AHL and their membrane composition, as both are Gram negative bacteria. DBHB is an anti-biofilm compound, but its activity is not permanent. If the molecule is diluted or removed, the biofilm is formed and the biovolume is not significantly different from the control ($p > 0.05$). To inhibit biofilm formation, DBHB has to be in contact with the bacteria.
were also tested and compared (Table 2).

±

sp. 4M6 and produce AHL (Pseudoalteromonas sp. 5M6) and a marine Gram positive bacterium (Bacillus sp. 4J6) were also tested and compared

viability, a staining with syto9® and sytoxRed® was performed.

Observation of Paracoccus sp. 4M6 biofilms (syto9®) with addition of DBHB at four concentrations (2, 8, 16 and 32 µM) in the medium growth flow, biovolumes (µm³/µm²) and biofilms percentages (%) are presented; the ± represents the standard deviation.

In order to determine the spectrum of activity of DBHB, a marine bacterium which did not produce AHL (Pseudoalteromonas sp. 5M6) and a marine Gram positive bacterium (Bacillus sp. 4J6) were also tested and compared (Table 2).

Table 2. Biovolumes of Bacillus sp. 4J6 and Pseudoalteromonas sp. 5M6 biofilms calculated for the control condition and after addition of DBHB at 16 and 32 µM in the growth medium flow; the ± represents the standard deviation.

| Bacterial strains | Control 16 µM DBHB | 32 µM DBHB |
|-------------------|---------------------|------------|
| 4J6 biovolumes (µm³/µm²) | 18.9 ± 1.8 | 21.5 ± 1.9 | 22.5 ± 1.7 |
| 5M6 biovolumes (µm³/µm²) | 22.0 ± 1.5 | 21.9 ± 1.3 | 22.1 ± 1.8 |
| ANOVA | - | - |
| TUKEY | - | - |

- no statistical difference, p > 0.05.

The addition of DBHB in the growth medium flow at 16 and 32 µM did not significantly disturb the biofilm formation of Bacillus sp. 4J6 and Pseudoalteromonas sp. 5M6. Biovolumes of Bacillus sp. 4J6 and Pseudoalteromonas sp. 5M6 biofilms were not different from the control condition (about 20 µm³/µm² for both bacteria with or without addition of DBHB). In this study, the molecule DBHB seemed to have a specific action on Gram negative bacteria which produce AHL, but more strains should be tested to confirm this result. As it was described that Gram negative bacteria are dominant in the marine environment, it is interesting to have an antibiofilm property on dominant bacteria in seawater [36].

2.4.2. Biofilm Degradation

The second way to evaluate the ability of DBHB to interfere with the biofilm’s behavior was to observe the impact of the compound on a biofilm already formed. DBHB was tested on two strains: Paracoccus sp. 4M6 and P. aeruginosa PAO1 (Table 3).

After a contact time of two hours between the bacterial biofilm and DBHB, the biofilm was observed. Biofilms that had already been formed were degraded by the addition of the DBHB in the channel flowcell. There was a factor of inhibition of 3 between the biovolume of the control biofilm (with biovolumes values of 12.9 ± 1.9 µm³/µm² for 4M6 and 14.6 ± 1.3 µm³/µm² for PAO1) and the biovolume after the contact with the DBHB (4.6 ± 1.1 µm³/µm² for 4M6 and of 4.5 ± 0.4 µm³/µm² for PAO1). These results were proved to be significantly different by a ANOVA (p < 0.01) test in comparison with the control.

To validate the activity of DBHB on biofilm formation without an interaction on the bacterial viability, a staining with syto9® and sytoxRed® was performed.
Table 3. Biovolumes of PAO1 and 4M6 biofilms tagged with syto9® and sytoxRed® to observe the impact of DBHB at 32 µM on the bacterial viability, the ± represents the standard deviation.

| Biovolumes (µm³/µm²)   | Paracoccus sp. 4M6 | Pseudomonas aeruginosa PAO1 |
|------------------------|-------------------|---------------------------|
|                        | Control          | 32 µM DBHB                | Control          | 32 µM DBHB       |
| Living cells (Syto9®)  | 12.7 ± 1.3       | 4.9 ± 0.1                 | 14.7 ± 1.0       | 4.3 ± 1.0       |
| Dead cells (SytoxRed®) | 0.2 ± 0.1        | 0.3 ± 0.1                 | 0.2 ± 0.02       | 0.2 ± 0.03      |
| Mortality percentage   | 0.98             | 0.94                      | 0.98             | 0.95            |
| ANOVA                  | -                | -                         | -                | -               |
| TUKEY                  | -                | -                         | -                | -               |

-: no statistical difference, p > 0.05.

The CLSM observations and the biovolume measurements showed that the addition of DBHB did not induce cells mortality within the biofilm. The percentage of altered cells tagged in red concomitantly with biovolume value (Table 1) indicated that DBHB did not have toxicity against those bacteria. Indeed, the biovolume of dead cells between the control and the biofilm after addition of DBHB were not significantly different (ANOVA and Tukey tests).

2.5. Anti-Quorum Sensing Activity

Continuing the demonstration of DBHB activity on biofilm formation, another test using a biosensor, E. coli pSB401, was performed in order to determine further biological properties of the compound on bacterial communication. In presence of AHL, the plasmid pSB401 containing QS genes in E. coli induces a luminescence which was possible to quantify. If DBHB has an impact on QS genes (luxCDABE), the luminescence production would be inhibited by the biosensor E. coli pSB401.

The luminescence production controlled by the recognition between AHL and the receptor is presented Figure 6. An inhibition of the luminescence was observed when the biosensor was in contact with DBHB. The inhibition appeared to be proportional to the concentration of DBHB and the molecule did not alter E. coli growth. The control displayed a classical curve of recognition between AHL and the receptor inducing the luminescence. The addition of DBHB at 8 and 16 µM inhibited the luminescence and the maximum values were respectively 1.7 and 2 times lower than those of the control. This inhibition is statistically different from the control (ANOVA and Tukey, p < 0.01). When using kojic acid, which is already known as a quorum sensing inhibitor, the inhibition was 20 times lower, as compared to the control.

![Figure 6](image-url)
To validate the anti-quorum sensing property of DBHB and the absence of interaction of the compound with the luminescence production, another biosensor, *V. harveyi* JAF548, was used. For this second biosensor, the bioluminescence production is not dependent of AHL presence. If the luminescence produced by *V. harveyi* JAF548 is inhibited, the molecule would not interact on QS genes but on the luminescence biosynthesis. Figure 7 presents the ratio between the bioluminescence emission and the optical density of *V. harveyi* JAF548 for the control condition and after the addition of DBHB at doses of 8 and 16 µM.

![Figure 7](image_url)

**Figure 7.** Ratio of the bioluminescence production and the optical density at 600 nm of the biosensor *V. harveyi* JAF548 with and without addition of DBHB at 8 and 16 µM; rods represent the standard deviation.

The biosensor *V. harveyi* JAF548 validates the absence of activity of DBHB on the bioluminescence production, confirming the anti-quorum sensing property of DBHB. At the maximum bioluminescence (4 h of growth), the production was not significantly different between the control condition and the addition of DBHB (ANOVA, *p* > 0.05).

3. Discussion

3.1. Anti-Bacterial Activity

DBHB is not toxic toward the bacteria studied. The absence toxicity is an important factor in order to allow the utilization of a new compound in an antifouling coating to be used in the natural environment. Nowadays, in order to authorize a new biocide, the molecule must have numerous requirements in order to obtain a marketing authorization, such as an absence of toxicity against referent organisms, and not be bioaccumulable. In Europe, a regulation on biocidal products (RBP) is established for the validation and use of biocides [37].

3.2. Impact of DBHB on AHL Production

Data show that DBHB does not impact the AHL production by *P. aeruginosa* PAO1 and *Paracoccus* sp. 4M6. Some quorum sensing inhibitor can degrade or perturb the AHL synthesis. For example, some bacteria produce enzymes such as lactonase or acylase, which can degrade AHL [15]. Without AHL, the bacterial communication is inhibited. By targeting the QS, it is possible to inhibit bacterial adhesion or biofilm maturation, which are important steps in microfouling [38].
3.3. Anti-Adhesion Activity

Experiments realized to assess the potential interaction of the DBHB on the bacterial adhesion, showed that no significant difference from the control was obtained for the bacteria *Paracoccus* sp. 4M6 and *Pseudomonas aeruginosa* PAO1. DBHB cannot inhibit the bacterial adhesion.

Other experiments not presented in this study showed that DBHB did not modify bacterial motility of *P. aeruginosa* PAO1: swimming, swarming and twitching. For this motility processes, two kinds of organelles, pili and flagellum, were used. Studies published that these organelles are involved in bacterial adhesion and biofilm structuration [39,40]. These results confirmed that DBHB did not impact cellular mechanism allowing the bacterial adhesion.

3.4. Anti-Biofilm Activity

For both strains *Paracoccus* sp. 4M6 and *P. aeruginosa* PAO1, DBHB led to an interesting activity on biofilm formation and biofilm desintegration. The marine strain *Paracoccus* sp. 4M6 was more sensitive than the terrestrial bacterium *P. aeruginosa* PAO1. From tested conditions in this study, DBHB has a specific activity towards Gram negative bacteria which produce AHL.

Gram negative bacteria possesses a complex cell wall, the DBHB could target a specific compound of the Gram negative cell wall which is not present in Gram positive bacteria, such as a lipopolysaccharide; it could cross the bacteria wall by porin protein [41]. In Gram negative bacteria, the outer membrane protects bacteria against antibiotics, detergents or other molecules [42]. It is an interesting result for antifouling application because, in the marine environment, Gram negative bacteria are dominant [36].

DBHB seems also able to target bacterial strains which produce bacterial signal molecule such as AHL; it is possible that this molecule can interfere in quorum sensing.

DBHB was derived from natural bastadins isolated from the marine sponge *Ianthella basta*. It was not surprising to find antibiofilm compounds in marine sponge; these organisms need to produce chemical defenses for their survival [43], against predation or for space and competition for resources [44,45]. Two classes of marine sponge compounds were previously identified as antibiofilm molecules without interaction on the bacterial viability, which include terpenoids [46] and the pyrrole imidazoles [47]. Some of these compounds are presented in Table 4.

### Table 4. Antibiofilm compounds derived from marine sponges without killing bacteria or disrupting their growth.

| Classes               | Compounds            | Activity                              | References          |
|-----------------------|----------------------|---------------------------------------|---------------------|
| Terpenoids            | Ageloxime-D         | antibiofilm, antimacrofouling          | [17,46]             |
|                       | Manoalide           | antibiotic, anti-inflammatory, anti-quorum sensing | [40]               |
| Pyrrole-imidazole     | Oroidin             | antibiofilm, antifouling               | [49–51]            |
| alkaloids             | Sceptrin            | antifouling, anti-quorum sensing       | [52]               |
|                       | Bromoageliferin     | anti-adhesion, antibiofilm             | [47,53]            |

Natural compounds which have this type of activity are a source of numerous analogues with biological activities. In this study, the bastadin analogue DBHB was found to be a promising antifouling compound, because this molecule can inhibit biofilm formation without interacting with bacterial growth. DBHB has a simple structure and can be easily synthesized.

DBHB is not the first compound exhibiting bromine atoms that show antibiofilm properties. Other studies have confirmed the antibiofilm property of brominated compounds, for example, in 2008 Huigens et al. showed that the increase of brominated groups in a given molecular structure induced a stronger antibiofilm activity [54]. In another study of Andjouh et al. in 2016, the authors demonstrated the importance of bromine groups for the antifouling performance [55].

An example to consider is the compound oroidin, isolated from the sponge *Agelas oroides*. The molecule has been described as an inhibitor of bacterial adhesion [56]. Forte et al. in 2009,
showed that oroidin can inhibit *P. aeruginosa* biofilm with IC$_{50}$ of 190 µM. Structurally, oroidin and DBHB are similar, with two aromatic cycles and bromines. By comparing, IC$_{50}$ of these two molecules on the inhibition of *P. aeruginosa* biofilms, DBHB appeared to be twenty times more active than oroidin [57].

Another explanation of DBHB effect could be an interaction with an intracellular pathway, such as with the C-di-GMP. This intracellular messenger is involved in biofilm formation and dispersion. In *P. aeruginosa*, the level of C-di-GMP induced the extracellular matrix synthesis by the activation of *pel* and *psl* genes [58].

3.5. Anti-Quorum Sensing Activity

The study with the biosensor *E. coli* pSB401 demonstrated an anti-QS property of DBHB. The addition of DBHB induces a reduction of the luminescence production; however, this inhibition is not similar to the activity of kojic acid. Kojic acid is described as an inhibitor of QS, which is explained by chemical similarities between AHL and kojic acid. The latter compound was identified as an AHL analogue. Kojic acid can bind to QS receptors such as RhlR, LasR or LuxR and block the communication system [59]. The difference on the inhibition of the bacterial communication by DBHB or acid kojic could be explained by the mode of action of the molecule. It is known that kojic acid possesses a direct activity on QS, as the molecule is an antagonist of AHL, while DBHB could interact on a step of regulation in the QS, which does not induce a total inhibition of the luminescence produced by the biosensor. With these results, we can hypothesize that DBHB does not interact in the same way as kojic acid does on the AHL receptor.

The second biosensor, *V. harveyi* JAF548 confirmed the activity of DBHB on QS and not on the bioluminescence production. Indeed, the strain *V. harveyi* JAF548 produces bioluminescence which is not under the control of QS. The absence of inhibition of bioluminescence validated the interaction of DBHB with QS [60].

Five steps of QS inhibitions were described, (i) compounds can interact on the biosynthesis of AHL; (ii) compounds can degrade QS signal molecules; (iii) compounds can inhibit AHL efflux protein; (iv) compounds can interact on transcriptional activator or (v) compounds can be considered as QS analogues [61]. It is possible that DBHB can interact on one of four of the last steps described.

DBHB is not the first derivative of compounds from sponges which presents an activity on QS. Another group of marine-sponge-derived compounds was demonstrated to be quorum sensing inhibitors: the manoalide family (Table 2). Extracts from the marine sponge *Luffariella variabilis* were shown to act as quorum sensing inhibitors [62], but the site of action has never been described before for these molecules. It is difficult to define precisely the action of a molecule on the QS because a lot of interactions are possible in the cell. Until now, some natural compounds were described as quorum sensing inhibitors; most compounds identified were able to inhibit the QS receptor such as manoalide or sceptrin type compounds [59]. To find new anti-quorum sensing compounds is an interesting way to disturb microbial communication linked in microfouling [63]. Some studies explained that they are links between QS and biofilm formation, but other studies described that too many pathways are involved to show a direct effect of QS on biofilm [64]. With genomic tools, it will be possible to determine the way of action of the molecule.

DBHB has already been described as an antifouling compound; the molecule can inhibit macrofoulers colonization. Its activity on an enzyme, the phenoloxydase, which occurs in the mussel adhesion, was described (IC$_{50}$ = 0.8 µM) [28]. The compound was also identified as an inhibitor of the barnacle adhesion (10 µM) [29]. In this study, DBHB is able to inhibit bacterial biofilm maturation (PAO1: IC$_{50}$ = 10.2 µM and 4M6: IC$_{50}$ < 2 µM) without toxicity, a possible explanation is that DBHB can interact on bacterial communication. Comparatively to macrofoulers, Gram-negative bacteria were less sensitive than mussels at DBHB, the IC$_{50}$ to inhibit biofilm maturation is 2 at 10 times higher than the enzymatic activity. All these data make DBHB a promising antifouling agent.
4. Materials and Methods

All chemical products were purchased from Sigma Aldrich except the syto9® which was purchased in Fisher scientific. DBHB had been synthesized at the Institute of pharmaceutical biology and biotechnology (Heinrich-Heine-University Düsseldorf, Germany) [27].

4.1. Bacterial Strains and Growth Conditions

Marine bacteria selected for this study were Paracoccus sp. 4M6 (Gram negative, 1 OD = \(4.8 \times 10^7\) CFU/mL), Pseudoalteromonas sp. 5M6 (Gram negative, 1 OD = \(3.5 \times 10^7\) CFU/mL) and Bacillus sp. 4J6 (Gram positive, 1 OD = \(4.2 \times 10^7\) CFU/mL). These marine bacteria were isolated from glass slides immersed for 6 h in the Morbihan gulf, France [65]. Paracoccus sp. 4M6 was selected for its ability to produce bacterial signal molecules: acyl homoserine lactone (AHL). In contrast, Pseudoalteromonas sp. 5M6 was chosen as a model species as it does not produce any AHL. Finally, Bacillus sp. 4J6 was selected as a representative of Gram positive bacteria. The three strains were maintained and cultivated at 20 °C using Zobell media (ASW 30 g/L, Tryptone 4 g/L, Yeast extract 1 g/L). Bacterial cultures were incubated at \(10^6\) UFC/mL for 48 h under agitation.

The biosensor Vibrio harveyi JAF548 was used to validate the absence of activity of DBHB on bioluminescence production [66]. The strain was grown in MB medium (37.5 g/L) supplemented with kanamycin at 30 μg/mL.

Two terrestrial bacterial strains were also selected. Pseudomonas aeruginosa PAO1 to evaluate antibiofilm activity (1 OD = \(2.7 \times 10^8\) CFU/mL), because it is a bacterium well known for biofilm formation and communication processes. The second terrestrial bacterium is the biosensor Escherichia coli pSB401 used to describe the anti-quorum sensing property. Terrestrial strains were maintained and cultivated at 37 °C in Luria Bertani, LB (NaCl 10 g/L, Tryptone 10 g/L, Yeast extract 5 g/L) for 24 h under agitation.

All experiments (anti-adhesion, anti-biofilm activities) were realized using Paracoccus sp. 4M6 and P. aeruginosa PAO1, however the remaining marine bacteria (Pseudoalteromonas sp. 5M6, Bacillus sp. 4J6, V. harveyi JAF548 and E. coli pSB401) were used to complete the activity of the DBHB on different bacteria.

4.2. Impact of DBHB on AHL Production

AHL was extracted and identified in four strains: Pseudomonas aeruginosa PAO1, Paracoccus sp. 4M6, Bacillus sp. 4J6 and Pseudoalteromonas sp. 5M6.

AHL extraction and identification processes were performed using the methodology described by Morin et al. [35]. The effect of DBHB was evaluated at three concentrations (16, 32 and 80 μM). In cultures with and without DBHB, AHL were extracted with dichloromethane in bacterial supernatants. The organic phase was dried and AHL were suspended with acetonitrile. Extracts were injected in LC–MS (microTOF-QII, Bruker, Billerica, MA, USA) and AHL were identified by analysis of the retention time and spectrum obtained [67]. All extractions were realized in triplicate.

4.3. Anti-Bacterial Activity

DBHB anti-bacterial activity was evaluated at six concentrations: 0.02, 0.08, 0.2, 2, 16 and 80 μM. Bioassays were run in twelve replicates. The screening was performed on Paracoccus sp. 4M6, Pseudoalteromonas sp. 5M6 and Pseudomonas aeruginosa PAO1. Bacterial cultures of 24 h for PAO1 and 48 h for 4M6 and 5M6 were diluted at \(10^6\) CFU/mL and added the microplate containing DBHB to a final volume of 200 μL in a 96-well microplate in polypropylene. DBHB was added in the first step to allow the evaporation of the solvent (methanol). A positive control consisted of bacteria without DBHB and for the negative control DCOIT (4,5-Dichloro-2-octyl-4-isothiazolin-3-one) was added at 0.1 μg/mL. Microplates were incubated at 20 °C for 48 h for 4M6 and 5M6, and at 37 °C for 24 h for PAO1.
After the incubation the optical density at 600 nm was measured to determine the impact of DBHB on bacterial growth. By a comparison to the control, a bacterial inhibition percentage was determined.

4.4. Anti-Adhesion Activities

In order to evaluate the anti-adhesion properties of DBHB, experiments were realized in a flowcell [30]. The flowcell was prepared by sticking a microscope coverslip 24 × 50 mm² (VWR) slide which is the support of adhesion. The flowcell is presented on the Figure 8.

The system was sterilized by a flow of bleach (1.5%) for 24 h at 430 µL/min. Then a flow of minimum medium, a saline solution (NaCl 9 g/L) for the terrestrial bacterium: *Pseudomonas aeruginosa* PAO1 or an artificial sea water (ASW 30 g/L) for marine bacteria (*Paracoccus* sp. 4M6, *Pseudoalteromonas* sp. 5M6 and *Bacillus* sp. 4J6) was activated to clean and prepare the system for the bacterial injection at 150 µL/min.

The bacterial solution was prepared from a bacterial culture which was inoculated overnight. A dilution of the bacterial suspension was realized to inject bacteria at 10⁶ CFU/mL in minimum medium at 16 µM due to a limitation of DBHB quantity available at the time. Also, to evaluate DBHB activity on bacterial adhesion, the molecule was used to prepare the support of adhesion. DBHB was added at 16 µM in the minimal medium flow. It was used to clean the surface during two h at 150 µL/min (Condition 2 in Table 5). A condition without addition of DBHB was used as a control.

After two hours of adhesion, the flow at 150 µL/min was activated during 30 min with the aim to remove free bacteria. Adhered bacteria were observed with Syto9® nucleic acid stain at 5 µM (λ<sub>excitation</sub> = 488 nm, λ<sub>emission</sub> = 498–540 nm). The bacterial adhesion was observed with confocal laser scanning microscopy (Zeiss, LSM 710, Oberkochen, Germany) by using a 40× oil immersion objective. For every studied parameter, a triplicate was realized (27 observations per condition). The overlap percentage was determined with a JAVA program established in the lab. This program determines a percentage between black and green pixels.
### 4.5. Anti-Biofilm Activities

In order to evaluate the anti-biofilm property of DBHB on three marine bacteria (*Paracoccus* sp. 4M6, *Pseudoalteromonas* sp. 5M6 and *Bacillus* sp. 4J6) and on one terrestrial bacterium *P. aeruginosa PAO1*, a test in a flowcell was performed. To observe the biofilm formation, a flow of medium growth at 120 µL/min (LB for *P. aeruginosa PAO1* or Zobell for *Paracoccus* sp. 4M6, *Pseudoalteromonas* sp. 5M6 and *Bacillus* sp. 4J6) was activated during 24 h for *P. aeruginosa PAO1* or 48 h for marine bacteria to allow the biofilm formation under dynamic condition. Then, bacteria were stained with the same procedure as the adhesion part with syto9® and the biofilm was observed by confocal laser scanning microscopy. The biovolume and the thickness of biofilms were determined with the COMSTAT program [68].

To highlight a potential anti-biofilm property of DBHB, two tests were realized. Firstly, DBHB activity on the biofilm formation was assessed. In this case, DBHB was added to the growth medium flow at 2, 8, 16 and 32 µM. Secondly, DBHB activity on the biofilm degradation was examined. For this second test, DBHB in solution at 16 µM in growth medium was injected in a channel with a formed biofilm. DBHB was left in contact with the biofilm for 2 h and the flow was activated to clean the channel before CLSM observations. The inhibition factor was calculated by a ratio between the percentage of biofilm between the control (bacteria without DBHB, 100%) and the percentage of biofilm obtained by the addition of the compound. For every concentration, a triplicate is realized (27 observations per condition).

The bacterial viability was measured using a double tagging by syto9® nucleic acid stain and sytox red® nucleic acid stain (λexcitation = 633 nm, λemission = 645–695 nm). The viability percentage was determined from biofilm biovolume.

### 4.6. Anti-Quorum Sensing Activities

In order to measure and characterize DBHB potential effect on QS, a biosensor was used. This biosensor constituted of a strain of *Escherichia coli* containing the plasmid pSB401. This plasmid harbors a luxCDABE gene which induces a luminescence when the promotor is activated in presence of AHL, the C6-HSL at 500 ng/mL. The selection of *E. coli* containing the plasmid pSB401 was controlled by a resistant gene at the tetracycline (10 µg/mL).

The test was carried out using a polypropylene 96-well black microplate. C6-HSL was used at 500 ng/mL to activate the biosensor; acetonitrile was used as carrier solvent for the AHL and was eliminated by evaporation under a laminar flow hood for 30 min. The 96-well was inoculated with *E. coli* pSB401 at 10⁶ UFC/mL in LB media with tetracycline at 10 µg/mL. DBHB was added at 8 and 16 µM to obtain a final volume of 200 µL per well. The 96-well microplate was incubated at 30 °C under agitation for 8 h. Every hour the Optical Density at 600 nm and the luminescence were measured (TECAN, infinite M200pro). A known inhibitor of quorum sensing, kojic acid, was also tested at 35 µM [59].

In order to validate the absence of bioluminescence production inhibition, another biosensor was used: *Vibrio harveyi* JAF548 was mutated on luxOD47E, the protein LuxO was maintained in...
a phosphorylated stage which allows continuous bioluminescence production. In this biosensor, the bioluminescence production is not controlled by quorum sensing and it is activated in continuous.

To evaluate the activity of DBHB, the bioluminescence emission reported to the optical density was measured every hour for twelve hours. A total of 100 μL of AB medium [69] containing the kanamycin at 30 μg/mL, Vibrio harveyi JAF548 at 10⁶ UFC/mL and DBHB evaluated at 2 and 8 μM were dropped. The experiment was performed in polypropylene 96-well black microplate.

All experiments were run in six replicates.

4.7. Statistical Analysis of Data

The means are reported ± standard deviation (SD). The effect of the compound DBHB on bacterial adhesion, biofilm formation, biofilm degradation, bacterial viability and bacterial communication were tested using the one-factor analysis variance (ANOVA) and the Tukey test (HSD). The level of significance was set to \( p < 0.01 \). Values are means ± standard deviation.

5. Conclusions

It was already shown that bacterial signal molecules such as AHL are involved in relationships between bacteria and other organisms, including microalgae [70] and macroorganisms (Ulva spores, Balanus cyprids) [71]. Targeting the bacterial communication system is an interesting strategy to inhibit, first, the microfouling by disruption of bacteria and diatom biofilms and, second, the macrofouling by the reduction of macrofoulers attachment. Dibromohemibastadin is a promising antifouling compound and we can now describe a first approach of this compound for the inhibition of microfouling.

In order to understand the way of action for this compound, a study on biofilm formation and quorum sensing was realized. DBHB was able to inhibit biofilm formation without acting on the adhesion step. The biosensor used, a modified E. coli strain, highlighted an interaction of DBHB with bacterial communication. The use of natural or analogue compounds, which have an impact on quorum sensing and biofilm formation, could be an interesting strategy to inhibit microfouling and macrofouling. In addition, the absence of toxicity of DBHB on bacteria is an important factor to consider that would allow DBHB to be applied as an AF agent. To evaluate the activity of DBHB on biofouling, it will be interesting to integrate the compound in a coating and immerse it in a natural environment.

Acknowledgments: The authors acknowledge the financial support for the Ph.D., the région Bretagne (ARED LACTOPAI) and the Université de Bretagne occidentale. Authors are gratefully acknowledged to Karen Tait from the Plymouth Marine Laboratory for the study with the strain E. coli pSB401 and to Ana Otero from the University of Santiago de Compostela for the strain V. harveyi JAF548. Authors want also to acknowledge Peter Proksch to allow the study on DBHB by the synthesis of the molecule.

Author Contributions: T.L.N. performed the experiments; F.F., K.R. and I.L. conceived and designed the experiments; I.L., F.F., K.R. and C.H. analyzed the data, K.T., H.N. and P.P. contributed to biosensor strain and DBHB providing; T.L.N. wrote the paper.

Conflicts of Interest: The authors declare no conflict of interest.

References

1. Wahl, M. Marine epibiosis. I. Fouling and antifouling: Some basic aspects. Mar. Ecol. Prog. Ser. 1989, 58, 175–189. [CrossRef]
2. Rubio, C. Compréhension des Mécanismes D’adhésion des Biofilms en Milieu Marin en vue de la Conception de Nouveaux Moyens de Prévention; Université Paris 6: Paris, France, 2002. (In French)
3. Qian, P.-Y.; Lau, S.C.K.; Dahms, H.-U.; Dobretsov, S.; Harder, T. Marine biofilms as mediators of colonization by marine macroorganisms: Implications for antifouling and aquaculture. Mar. Biotechnol. 2007, 9, 399–410. [CrossRef] [PubMed]
4. Callow, J.A.; Callow, M.E. Trends in the development of environmentally friendly fouling-resistant marine coatings. Nat. Commun. 2011, 2, 244. [CrossRef] [PubMed]
5. Tsuneda, S.; Aikawa, H.; Hayashi, H.; Yuasa, A.; Hirata, A. Extracellular polymeric substances responsible for bacterial adhesion onto solid surface. *FEMS Microbiol. Lett.* 2003, 223, 287–292. [CrossRef]  
6. Waters, C.M.; Bassler, B.L. Quorum sensing: Cell-to-Cell Communication in Bacteria. *Annu. Rev. Cell Dev. Biol.* 2005, 21, 319–346. [CrossRef] [PubMed]  
7. Waters, C.M.; Lu, W.; Rabinowitz, J.D.; Bassler, B.L. Quorum sensing controls biofilm formation in *Vibrio cholerae* through modulation of cyclic Di-GMP levels and repression of vpsT. *J. Bacteriol.* 2008, 190, 2527–2536. [CrossRef] [PubMed]  
8. Schaefer, A.L.; Hanzelka, B.L.; Eberhard, A.; Greenberg, E.P. Quorum sensing in *Vibrio fischeri*: Probing autoinducer-LuxR interactions with autoinducer analogs. *J. Bacteriol.* 1996, 178, 2897–2901. [CrossRef] [PubMed]  
9. Cook, L.C.; Federle, M.J. Peptide pheromone signaling in *Streptococcus* and *Enterococcus*. *FEMS Microbiol. Rev.* 2014, 38, 473–492. [CrossRef] [PubMed]  
10. Davies, D.G.; Parsek, M.R.; Pearson, J.P.; Iglewski, B.H.; Costerton, J.W.; Greenberg, E.P. The involvement of Cell-to-Cell signals in the development of a bacterial biofilm. *Science* 1998, 280, 295–298. [CrossRef] [PubMed]  
11. Miller, M.B.; Bassler, B.L. Quorum sensing in bacteria. *Annu. Rev. Microbiol.* 2001, 55, 165–199. [CrossRef] [PubMed]  
12. Manefield, M.; Rasmussen, T.B.; Henzter, M.; Andersen, J.B.; Steinberg, P.; Kjelleberg, S.; Givskov, M. Halogenated furanones inhibit quorum sensing through accelerated LuxR turnover. *Microbiology* 2002, 148, 1119–1127. [CrossRef] [PubMed]  
13. Kociolek, M.G. Quorum-sensing inhibitors and biofilms. *Anti-Infect. Agents Med. Chem.* 2009, 8, 315–326. [CrossRef]  
14. Manefield, M.; de Nys, R.; Naresh, K.; Roger, R.; Givskov, M.; Peter, S.; Kjelleberg, S. Evidence that halogenated furanones from *Delisea pulchra* inhibit acylated homoserine lactone (AHL)-mediated gene expression by displacing the AHL signal from its receptor protein. *Microbiology* 1999, 145, 283–291. [CrossRef] [PubMed]  
15. Dong, Y.-H.; Zhang, L.-H. Quorum sensing and quorum-quenching enzymes. *J. Microbiol.* 2005, 43, 101–109. [PubMed]  
16. Yada, S.; Kamalesh, B.; Sonwane, S.; Guptha, I.; Swetha, R.K. Quorum sensing inhibition, relevance to periodontics. *J. Int. Oral Health* 2015, 7, 67–69. [PubMed]  
17. Fusetani, N. Antifouling marine natural products. *Nat. Prod. Rep.* 2011, 28, 400–410. [CrossRef] [PubMed]  
18. Qian, P.-Y.; Li, Z.; Xu, Y.; Li, Y.; Fusetani, N. Mini-review: Marine natural products and their synthetic analogs as antifouling compounds: 2009–2014. *Biofouling* 2015, 31, 101–122. [CrossRef] [PubMed]  
19. Sjögren, M.; Johnson, A.-L.; Hedner, E.; Dahlström, M.; Göransson, U.; Shirani, H.; Bergman, J.; Jonsson, P.R.; Bohlin, L. Antifouling activity of synthesized peptide analogs of the sponge metabolite barettin. *Peptides* 2006, 27, 2058–2064. [CrossRef] [PubMed]  
20. Perdicaris, S.; Vlachogianni, T.; Valavanidis, A. Bioactive natural substances from marine sponges: New developments and prospects for future pharmaceuticals. *Nat. Prod. Chem. Res.* 2013, 1, 2329–6836. [CrossRef]  
21. Laport, M.S.; Santos, O.C.S.; Muricy, G. Marine sponges: Potential sources of new antimicrobial drugs. *Curr. Pharm. Biotechnol.* 2009, 10, 86–105. [CrossRef] [PubMed]  
22. Thomas, T.R.A.; Kavlekar, D.P.; LokaBharathi, P.A. Marine drugs from sponge-microbe association—A review. *Mar. Drugs* 2010, 8, 1417–1468. [CrossRef] [PubMed]  
23. Wijffels, R.H. Potential of sponges and microalgae for marine biotechnology. *Trends Biotechnol.* 2008, 26, 26–31. [CrossRef] [PubMed]  
24. Sjögren, M.; Dahlström, M.; Hedner, E.; Jonsson, P.R.; Vik, A.; Gundersen, L.-L.; Bohlin, L. Antifouling activity of the sponge metabolite agarasin D and synthesized analogs on Balanus improvisus. *Biofouling* 2008, 24, 251–258. [CrossRef] [PubMed]  
25. Nir, S.; Reches, M. Bio-inspired antifouling approaches: The quest towards non-toxic and non-biocidal materials. *Curr. Opin. Biotechnol.* 2016, 39, 48–55. [CrossRef] [PubMed]  
26. Müller, W.E.G.; Wang, X.; Proksch, P.; Perry, C.C.; Osinga, R.; Gardères, J.; Schröder, H.C. Principles of biofouling protection in marine sponges: A model for the design of novel biomimetic and bio-inspired coatings in the marine environment? *Mar. Biotechnol.* 2013, 15, 375–398. [CrossRef] [PubMed]
27. Bayer, M.; Hellio, C.; Maréchal, J.-P.; Frank, W.; Lin, W.; Weber, H.; Proksch, P. Antifouling bastadin congeners target mussel phenoloxidase and complex copper(II) ions. *Mar. Biotechnol.* 2011, 13, 1148–1158. [CrossRef] [PubMed]
28. Niemann, H.; Hagenow, J.; Chung, M.-Y.; Hellio, C.; Weber, H.; Proksch, P. SAR of sponge-inspired hemibastadin congeners inhibiting blue mussel phenoloxidase. *Mar. Drugs* 2015, 13, 3061–3071. [CrossRef] [PubMed]
29. Ortlepp, S.; Sjögren, M.; Dahlström, M.; Weber, H.; Ebel, R.; Edrada, R.; Thoms, C.; Schupp, P.; Bohlin, L.; Proksch, P. Antifouling activity of bromotyrosine-derived sponge metabolites and synthetic analogues. *Mar. Biotechnol.* 2007, 9, 776–785. [CrossRef] [PubMed]
30. Tolker-Nielsen, T.; Sternberg, C. Methods for studying biofilm formation: Flow cells and confocal laser scanning microscopy. In *Pseudomonas Methods and Protocols*; Filloux, A., Ramos, J.-L., Eds.; Methods in Molecular Biology; Springer: New York, NY, USA, 2014; pp. 615–629; ISBN 978-1-4939-0472-3.
31. Grasland, B. Etude du Rôle des Interactions Energétiques et des Communications Intercellulaires Dans la Formation de Biofilms Bactériens en Milieu Marin. Ph.D. Thesis, Université de Bretagne-Sud, Lorient, France, 2002.
32. Jacobson, A.H.; Willingham, G.L. Sea-nine antifoulant: An environmentally acceptable alternative to organotin antifoulants. *Sci. Total Environ.* 2000, 258, 103–110. [CrossRef]
33. Yang, J.; Evans, B.A.; Rozen, D.E. Signal diffusion and the mitigation of social exploitation in pneumococcal competence signalling. *Proc. Biol. Sci.* 2010, 277, 2991–2999. [CrossRef] [PubMed]
34. Monnet, V.; Juillard, V.; Gardan, R. Peptide conversations in Gram-positive bacteria. *Crit. Rev. Microbiol.* 2016, 42, 339–351. [CrossRef] [PubMed]
35. Morin, D.; Grasland, B.; Vallée-Réhel, K.; Dufau, C.; Haras, D. On-line high-performance liquid chromatography-mass spectrometric detection and quantification of N-acylhomoserine lactones, quorum sensing signal molecules, in the presence of biological matrices. *J. Chromatogr. A* 2003, 1002, 79–92. [CrossRef]
36. Zhang, Y.; Jiao, N.; Cottrell, M.T.; Kirchman, D.L. Contribution of major bacterial groups to bacterial biomass production along a salinity gradient in the South China Sea. *Aquat. Microb. Ecol.* 2006, 43, 233–241. [CrossRef]
37. Hommen, U.; Baveco, J.M.; Galic, N.; van den Brink, P.J. Potential application of ecological models in the European environmental risk assessment of chemicals I: Review of protection goals in EU directives and regulations. *Integr. Environ. Assess. Manag.* 2010, 6, 325–337. [CrossRef] [PubMed]
38. Ponnusamy, K.; Paul, D.; Kweon, J.H. Inhibition of quorum sensing mechanism and aeromonas hydrophila biofilm formation by vanillin. *Environ. Eng. Sci.* 2009, 26, 1359–1363. [CrossRef]
39. O’Toole, G.A.; Kolter, R. Flagellar and twitching motility are necessary for Pseudomonas aeruginosa biofilm formation by vanillin. *Mar. Drugs* 2011, 13, 1148–1158. [CrossRef] [PubMed]
40. Toutain, C.M.; Caiazza, N.C.; Zegans, M.E.; O’Toole, G.A. Roles for flagellar stators in biofilm formation by Pseudomonas aeruginosa. *Mar. Drugs* 2017, 15, 222. [CrossRef]
41. DiRienzo, J.M.; Nakamura, K.; Inouye, M. The outer membrane proteins of gram-negative bacteria: Biosynthesis, assembly, and functions. *Annu. Rev. Biochem.* 1978, 47, 481–532. [CrossRef] [PubMed]
42. Nikaido, H. Multidrug efflux pumps of gram-negative bacteria. *J. Bacteriol.* 1996, 178, 5853–5859. [CrossRef] [PubMed]
43. Braekman, J.C.; Daloze, D. Chemical defence in sponges. *Pure Appl. Chem.* 2009, 58, 357–364. [CrossRef]
44. Engel, S.; Pawlik, J.R. Allelopathic activities of sponge extracts. *Mar. Ecol. Prog. Ser.* 2000, 207, 273–281. [CrossRef]
45. Assmann, M.; Lichte, E.; Pawlik, J.R.; Kck, M. Chemical defenses of the Caribbean sponges Agelas wiedenmayeri and Agelas conifera. *Mar. Ecol. Prog. Ser.* 2000, 207, 255–262. [CrossRef]
46. Hertiani, T.; Edrada-Ebel, R.; Ortlepp, S.; van Soest, R.W.M.; de Voogd, N.J.; Wray, V.; Hentschel, U.; Kozytska, S.; Müller, W.E.G.; Proksch, P. From anti-fouling to biofilm inhibition: New cytotoxic secondary metabolites from two Indonesian Agelas sponges. *Bioorg. Med. Chem.* 2016, 24, 1297–1311. [CrossRef] [PubMed]
47. Huigens, R.W.; Richards, J.J.; Parise, G.; Ballard, T.E.; Zeng, W.; Deora, R.; Melander, C. Inhibition of *Pseudomonas aeruginosa* biofilm formation with bromoageliferin analogues. *J. Am. Chem. Soc.* 2007, 129, 6966–6967. [CrossRef] [PubMed]
48. Ebada, S.S.; Lin, W.; Proksch, P. Bioactive sesterterpenes and triterpenes from marine sponges: Occurrence and pharmacological significance. *Mar. Drugs* 2010, 8, 313–346. [CrossRef] [PubMed]
49. Melander, C.; Moeller, P.D.R.; Ballard, T.E.; Richards, J.J.; Huigens, R.W., III; Cavanagh, J. Evaluation of dihydromororidin as an antifouling additive in marine paint. Int. Biodeterior. Biodegrad. 2009, 63, 529–532. [CrossRef] [PubMed]

50. Trepas, R.; Cervin, G.; Hellio, C.; Pavia, H.; Stensen, W.; Stensvåg, K.; Svendsen, J.-S.; Haug, T.; Svenson, J. Antifouling compounds from the sub-arctic ascidian synoicum pulmonaria: Synovoxazolidinones A and C, pulmonarins A and B, and synthetic analogues. J. Nat. Prod. 2014, 77, 2105–2113. [CrossRef] [PubMed]

51. Ballard, T.E.; Richards, J.J.; Wolfe, A.L.; Melander, C. Synthesis and antibiofilm activity of a second-generation reverse-Amide oroidin library: A structure-activity relationship study. Chem. Eur. J. 2008, 14, 10745–10761. [CrossRef] [PubMed]

52. Chifiriuc, C.M.; Grumezescu, M.A.; Lazar, V. Quorum sensing inhibitors from the sea: Lessons from marine symbiotic relationships. Curr. Organ. Chem. 2014, 18, 823–839. [CrossRef]

53. Nithya, C.; Begum, M.F.; Pandian, S.K. Marine bacterial isolates inhibit biofilm formation and disrupt mature biofilms of Pseudomonas aeruginosa PAO1. Appl. Microbiol. Biotechnol. 2010, 88, 341–358. [CrossRef] [PubMed]

54. IIii, R.W.H.; Ma, L.; Gambino, C.; Moeller, P.D.R.; Basso, A.; Cavanagh, J.; Wozniak, D.J.; Melander, C. Control of bacterial biofilms with marine alkaloid derivatives. Mol. BioSyst. 2008, 4, 614–621. [CrossRef]

55. Andjouh, S.; Blache, Y. Screening of bromotyramine analogues as antifouling compounds against marine bacteria. Biofouling 2016, 32, 871–881. [CrossRef] [PubMed]

56. Richards, J.J.; Ballard, T.E.; Huigens, R.W.; Melander, C. Synthesis and screening of an oroidin library against Pseudomonas aeruginosa biofilms. Mar Drugs 2009, 7, 705–753. [CrossRef] [PubMed]

57. Ha, D.-G.; O’Toole, G.A. c-di-GMP and its effects on biofilm formation and dispersion: A Pseudomonas aeruginosa review. Microbiol. Spectr. 2015, 3. [CrossRef] [PubMed]

58. Dobretsov, S.; Teplitski, M.; Bayer, M.; Gunasekera, S.; Proksch, P.; Paul, V.J. Inhibition of marine biofouling by bacterial quorum sensing inhibitors. Biofouling 2011, 27, 893–905. [CrossRef] [PubMed]

59. Dobretsov, S.; Teplitski, M.; Bayer, M.; Gunasekera, S.; Proksch, P.; Paul, V.J. Inhibition of marine biofouling by bacterial quorum sensing inhibitors. Biofouling 2011, 27, 893–905. [CrossRef] [PubMed]

60. Anetzberger, C.; Pirch, T.; Jung, K. Heterogeneity in quorum sensing-regulated bioluminescence of Vibrio harveyi. Mol. Microbiol. 2009, 73, 267–277. [CrossRef] [PubMed]

61. Kalia, V.C.; Kumar, P.; Pandian, S.T.K.; Sharma, P. Biofouling control by quorum quenching. In Springer Handbook of Marine Biotechnology; Prof, S.-K.K., Ed.; Springer: Berlin/Heidelberg, Germany, 2015; pp. 431–440. ISBN 978-3-642-53970-1.

62. Skinderoesoe, M.E.; Ettinger-Epstein, P.; Rasmussen, T.B.; Bjarnsholt, T.; de Nys, R.; Givskov, M. Quorum sensing antagonism from marine organisms. Mar. Biotechnol. 2007, 10, 56–63. [CrossRef] [PubMed]

63. Dobretsov, S.; Teplitski, M.; Bayer, M.; Gunasekera, S.; Proksch, P.; Paul, V.J. Inhibition of marine biofouling by bacterial quorum sensing inhibitors. Biofouling 2011, 27, 893–905. [CrossRef] [PubMed]

64. Kievit, T.R.D.; Gillis, R.; Marx, S.; Brown, C.; Iglewski, B.H. Quorum-sensing genes in Pseudomonas aeruginosa biofilms: Their role and expression patterns. Appl. Environ. Microbiol. 2001, 67, 1865–1873. [CrossRef] [PubMed]

65. Grasland, B.; Mitalane, J.; Briandet, R.; Quemener, E.; Meylheuc, T.; Linossier, I.; Vallee-Rehel, K.; Haras, D. Bacterial biofilm in seawater: Cell surface properties of early-attached marine bacteria. Biofouling 2003, 19, 307–313. [CrossRef] [PubMed]

66. Freeman, J.A.; Bassler, B.L. A genetic analysis of the function of LuxO, a two-component response regulator involved in quorum sensing in Vibrio harveyi. Mol. Microbiol. 1999, 31, 665–677. [CrossRef] [PubMed]

67. Fekete, A.; Frommberger, M.; Rothballer, M.; Li, X.; Englmann, M.; Fekete, J.; Hartmann, A.; Eberl, L.; Schmitt-Kopplin, P. Identification of bacterial N-acylhomoserine lactones (AHLs) with a combination of ultra-performance liquid chromatography (UPLC), ultra-high-resolution mass spectrometry, and in-situ biosensors. Anal. Bioanal. Chem. 2007, 387, 455–467. [CrossRef] [PubMed]

68. Heydorn, A.; Nielsen, A.T.; Hentzer, M.; Sternberg, C.; Givskov, M.; Ersbøll, B.K.; Molin, S. Quantification of biofilm structures by the novel computer program comstat. Microbiology 2000, 146, 2395–2407. [CrossRef] [PubMed]

69. Greenberg, E.P.; Hastings, J.W.; Ulitzur, S. Induction of luciferase synthesis in Beneckea harveyi by other marine bacteria. Arch. Microbiol. 1979, 120, 87–91. [CrossRef]
70. Yang, C.; Fang, S.; Chen, D.; Wang, J.; Liu, F.; Xia, C. The possible role of bacterial signal molecules $N$-acyl homoserine lactones in the formation of diatom-biofilm (Cylindrotheca sp.). *Mar. Pollut. Bull.* **2016**, *107*, 118–124. [CrossRef] [PubMed]

71. Tait, K.; Havenhand, J. Investigating a possible role for the bacterial signal molecules $N$-acylhomoserine lactones in Balanus improvisus cyprid settlement. *Mol. Ecol.* **2013**, *22*, 2588–2602. [CrossRef] [PubMed]

© 2017 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (http://creativecommons.org/licenses/by/4.0/).