Psammaceratin A: A Cytotoxic Psammaplysin Dimer Featuring an Unprecedented (2Z,3Z)-2,3-Bis(aminomethylene)succinamide Backbone from the Red Sea Sponge Pseudoceratina arabica

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Abstract: Bioassay-guided partition of the extract of the Red Sea sponge Pseudoceratina arabica and HPLC purification of the active fraction gave a psammaplysin dimer, psammaceratin (1), along with psammaplysin A (2). The dimer comprises two units of psammaplysin A (2) connected via the terminal amines with an unprecedented (2Z,3Z)-2,3-bis(aminomethylene)succinamide moiety, and it represents the first dimer to be identified among the psammaplysin family. Data from 1D- and 2D-NMR and HRMS supported the chemical structures of the compounds. Psammaceratin A (1) and psammaplysin A (2) exhibited significant growth inhibition of HCT 116, HeLa, and MBA-MB-231 cells down to 3.1 µM.

Keywords: Red Sea sponge; Pseudoceratina arabica; marine alkaloids; psammaplysin dimer; psammaceratin A; psammaplysin A; cell lines’ growth inhibition
backbone, another dibrominated bromotyrosine-derived unit named moloka’iamine [2],
connected together via an amidic linkage [13]. Psammaplysins’ derivatives have been
associated with diverse pharmacological properties, such as cytotoxic [5,11,13], antimi-
gratory [17], antimalaria [8,10,14,15], antiviral [6], antifouling [14], antimicrobial [18],
antioxidant [16], and immunosuppressive [6] activities.

In a continuation of our survey of Verongid sponges in the Red Sea for bioactive
chemical entities [13,19–22], bioassay-guided fractionation of the cytotoxic fraction of
the methanolic extract of P. arabica afforded a novel symmetrical psammaplysin dimer,
psammaceratin A (1), and the known psammaplysin A (2) [4]. Psammaceratin A (1)
possesses an unprecedented moiety, \((2Z,3Z)\)-2,3-bis(aminomethylene)succinamide, linking
symmetrically through the terminal amino groups of two units of psammaplysin A with
C-21/C-21′ of the substituted 2,3-bis(aminomethylene)succinamide part of the molecule.

2. Results and Discussion

2.1. Structure of Psammaceratin A (1)

Psammaceratin A (1) (Figure 1) was obtained as an optically active \((\left[\alpha\right]_D^{25} = -59°)\)
compound. The existence of the ion peaks at \(m/z\) 1616.6, 1618.6, 1620.6, 1622.6, 1624.6, 1626.6,
1628.6, 1630.6, and 1632.6 (1:8:28:56:70:56:28:8:1, \([M + Na]^+\)) in the positive ESIMS of 1
supported the presence of eight bromine atoms. The molecular formula of 1 was established
as \(\text{C}_{48}\text{H}_{50}\text{Br}_8\text{N}_8\text{O}_{14}\) by positive HRESIMS (\(m/z\) 1624.6732, \(\text{C}_{48}\text{H}_{50}\text{Br}_4\text{Na}_8\text{O}_{14}\),
\([M + Na]^+\)) (Figure S1), suggesting 24 degrees of unsaturation. Its \(^{13}\text{C}\) NMR spectrum
exhibited 22 resonances, corresponding to 24 carbons, signifying the presence of a sym-
mmetrical dimer (Table 1). Investigation of the complete NMR spectra of 1 including \(^1\text{H}\)
(Figures S2 and S3), \(^{13}\text{C}\) (Figure S4), DEPT (Figure S5), \(^1\text{H}-^1\text{H COSY}\) (Figure S6), and
multiplicity-edited HSQC (Figure S7) experiments confirmed the existence of five struc-
tural subunits in 1, including two similar pairs (A and A′, B and B′) together with an
unprecedented moiety (fragment C) (Table 1 and Figure 2).

![Figure 1. Chemical structures of 1 and 2.](image-url)
Table 1. NMR data of 1 (600 MHz for $^1$H and 150 for $^{13}$C, CD$_3$OD).

| No.   | $\delta$ C | $\delta$ H | HMBC            | NOESY         |
|-------|------------|------------|-----------------|---------------|
| 1, 1' | 146.8, CH  | 7.13 (s)   | C-2/2', C-3/3', C-6/6' | H$_2$-11/11' |
| 2, 2' | 104.4, qC  |            |                  |               |
| 3, 3' | 149.9, qC  |            |                  |               |
| 4, 4' | 104.6, qC  |            |                  |               |
| 5a, 5a' | 38.3, CH$_2$ | 3.38 (d, 16.2) | C-3/3', C-4/4', C-6/6', C-7/7' | H-5b/5b', H-7/7' |
| 5b, 5b' | 3.05 (d, 16.2) |            |                  | H-5a/5a', H-5b/5b' |
| 6, 6' | 120.9, qC  |            |                  |               |
| 7, 7' | 80.5, CH   | 4.98 (s)   | C-6/6', C-8/8', C-9/9' | H-5a/5a', H-5b/5b' |
| 8, 8' | 158.8, qC  |            |                  |               |
| 9, 9' | 160.6, qC  |            |                  |               |
| 10, 10' | 38.1, CH$_2$ | 3.60 (t, 6.5) |                  |               |
| 11, 11' | 30.6, CH$_2$ | 2.12 (quin., 6.5) |                  | H-5b/5b', H-9/9', H-10/10', H$_2$-12' |
| 12, 12' | 72.2, CH$_2$ | 4.05 (t, 6.5) |                  | H$_2$-10/10', H$_2$-11/11' |
| 13, 13' | 153.4, qC  |            |                  |               |
| 14, 14' | 119.4, qC  |            |                  |               |
| 15, 15' | 134.8, CH  | 7.43 (s)   | C-13/13', C-14/14', C-16/16', C-18/18', C-19/19' | H$_2$-19/19' |
| 16, 16' | 138.1, qC  |            |                  |               |
| 17, 17' | 134.8, CH  | 7.43 (s)   | C-13/13', C-14/14', C-16/16', C-18/18', C-19/19' | H$_2$-19/19' |
| 18, 18' | 119.4, qC  |            |                  |               |
| 19, 19' | 36.1, CH$_2$ | 2.90 (t, 7.2) | C-15/15', C-16/16', C-17/17', C-20/20' | H$_2$-20/20' |
| 20, 20' | 52.6, CH$_2$ | 3.69 (t, 7.2) | C-16/16', C-19/19', C-21/21' | H$_2$-19/19' |
| 21, 21' | 157.1, CH  | 7.54 (s)   | C-20/20', C-22/22', C-23/23' |               |
| 22, 22' | 119.2, qC  |            |                  |               |
| 23, 23' | 171.1, qC  |            |                  |               |
| 24, 24' | 59.4, CH$_3$ | 3.64 (s)   |                  | C-3/3'        |

Figure 2. Structural subunits and significant COSY and HMBC of 1.

The first similar subunits (A and A') in 1 are assigned as 2,3,4,7,9-penta-substituted spirooxepinisoxazoline moieties. These assignments are supported by the $^1$H and $^{13}$C NMR resonances at $\delta$$_H$/C 7.13 (s)/146.8 (CH, C-1/1'), 104.4 (qC, C-2/2'), 1549.9 (qC, C-3/3'), 104.6 (qC, C-4/4'), 3.38 (d, $^{2}$J = 16.2 Hz) and 3.05 (d, $^{2}$J = 16.2 Hz)/38.3 (CH$_2$, C-5/5'), 120.9 (qC, C-6/6'), 4.98 (s)/80.5 (CH, C-7/7'), 158.8 (qC, C-8/8'), 160.7 (qC, C-9/9'), and 3.64 (s)/59.4 (CH$_3$, C-25/25'). These signals are characteristic of the presence of two similar moieties, namely, 2,3,4,7,9-penta-substituted spirooxepinisoxazoline [2–13]. The locations of the bromine atoms, OCH$_3$, and OH moieties on the spirooxepinisoxazoline moieties at C-2/2', C-4/4', C-3/3', and C-7/7', respectively, were supported by HMBC.
cross-peaks from H-1/1' to C-2/2', C-3/3', and C-6/6'; from H2-5/5' to C-3/3', C-4/4', and C-6/6'; from H-7/7' to C-6/6', C-8/8', and C-9/9'; and from H2-24/24' to C-3/3' (Table 1 and Figure 2). Furthermore, these HMBC correlations supported the assignment of the quaternary carbons in subunits A and A' and complete the unequivocal assignment of the first subunits (A and A') of 1 (Figure 2).

The second identical subunits of 1 (B and B') (Figure 2) are assigned as two N,N-disubstituted moloka'i amine moieties. Two spin–spin coupling systems from H2-10/10' to H2-12/12' and between H2-19/19' and H2-20/20' were traced from the COSY experiment. The δH/C 3.60 (t, J = 6.5 Hz)/38.1 (CH2, C-10/10'), 2.12 (quin., J = 6.5 Hz)/30.6 (CH2, C-11/11'), 4.05 (t, J = 6.5 Hz)/72.2 (CH2, C-12/12'), 153.4 (qC, C-13/13'), 191.4 (2 × qC, C-14/14' and C-18/18'), 7.43 (s)/134.8 (2 × CH, C-15/15' and C-17/17'), 138.1 (qC, C-16/16'), 2.90 (t, J = 7.2 Hz)/36.1 (CH2, C-19/19'), and 3.69 (t, J = 7.2 Hz)/52.6 (CH2, C-20/20') suggested the existence of two moloka'i amine moieties, substituted on their terminal amines [14,15]. There was a significant downfield shift of the 13C NMR signals of the methylene moieties C-19/19' and C-20/20' and the 1H NMR signals of H2-19/19' in 1 in comparison with psammaplysins A (2), which possesses a free terminal amine at C-20. The change in the NMR shifts of these signals is attributed to the substitution of the free amines at C-20/20' in 1 by a substituted succinimide moiety as discussed below (Table 2).

The HMBC correlations (Table 1 and Figure 2 and Figure S8) from H2-10/10' to C-11/11' and C-12/12'; from H2-11/11' to C-10/10' and C-12/12'; from H2-12/12' to C-10/10', C-11/11', and C-13/13' (δC 153.4); from H-15/15' and H-17/17' to C-13/13', C-14/14', C-18/18' (δC 119.4), and C-19/19' (δC 36.1); from H-21/21' to C-16/16' (δC 138.1), C-17/17', and C-20/20'; and, finally, from H2-20/20' to C-16/16' and C-19/19' secured the structure of the subunits B and B' of 1.

Table 2. Comparison of partial NMR data of 1 and 2 (CD3OD).

| Position (δC ppm) | Psammaplysins A (2) | Psammaceratin A (1) | ΔδH (ppm) | ΔδC (ppm) |
|------------------|----------------------|----------------------|-----------|-----------|
| 19/19'           | 31.8                 | 36.1                 | -0.03     | +5.3      |
| 20/20'           | 40.0                 | 52.6                 | +0.51     | +12.6     |

(*) Data from this study.

The connections between the subunits A and B and between A' and B' through the amicd linkages C-9−N and C-9′−N′ are reinforced by HMBC correlations from H2-10/10' (δH 3.60) to C-9'/9' (δC 160.7) and from H-7/7' (δH 4.72 (Table 1 and Figure 2 and Figure S8)). This assignment was established by HMBC correlations from H2-20/20' to C-21/21', from H-21/21' to C-20/20', from H-21/21' to C-22/22', and from H-21/21' to C-23/23', completing the assignment of fragment C.

The sum of the elements of the assigned subunits A, A', B, and B' was counted for C42H44Br8N6O12 and for 20 degrees of unsaturation. The remaining elements of C6H8N2O2 (Fragment C) were counted for the remaining four degrees of unsaturation in 1. These elements C6H8N2O2 are assigned as 2,3-bis(aminomethylene)succinimide (Figure 2) based on the remaining NMR resonances at δH/C (CH, 7.54/157.1, C-21/21'), 119.2 (qC, C-22/22'), and 171.1 (qC, C-23/23') (Table 1 and Figure 2 and Figure S8). The linkage of subunit C with structural parts B and B' through the terminal amines and C-21/21' (NH-C-21 and NH-C-21') is supported by the long-range COSY couplings (‘J) between H2-20/20' (δH 3.69) and H-21/21' (δH 7.54), as well by HMBC correlations from H2-20/20' (δH 3.69) to C-21/21' (δC 157.1) and from H-21/21' (δH 7.54) to C-20/20' (δC 52.6) (Table 1 and Figure 2 and Figure S8), completing the degrees of unsaturation and the planar structure of psammaceratin A.

The substitution of the terminal amines in 1 with 2,3-bis(aminomethylene)succinimide moiety caused a significant and expected 13C NMR downfield shift of the carbons of the ethylene moieties (C-19/19' and C-20/20') from δC 31.8 (C-19) and δC 40.0 (C-20) in psammaplysins A (2) to δC 36.1 (C-19/19') and δC 52.6 (C-20/20') (ΔδC = +5.3 and +12.6 ppm,
respectively) in 1 (Table 2). An additional significant and expected downfield shift of H$_{-20}$ from 3.18 ppm (in 2) to 3.69 ppm in 1 ($\Delta$δ$_{H} = +0.51$ ppm) was observed, confirming the effect of the substitution of the terminal amines in psammaplysin derivatives [11] (Table 2).

The $\Delta$21,22 and $\Delta$21',22' configurations of the 2,3-bis(aminomethylene)succinamide moiety were assigned as Z and Z based on the presence of the NOE correlations between H$_{-20}$/20 and H-21/21' in the NOESY experiment (Figure 3 and Figure S9). In an MM2 energy-minimized drawing of 1 (Figure 4), strong NOE correlations are expected between H$_{-20}$ and H-21 and between H-20' and H-21' (Figures 3 and 4). Accordingly, the Z configurations at $\Delta$21,22 and $\Delta$21',22' are confirmed.

Figure 3. Significant NOESY correlations at the Z-configured $\Delta$21,22 and $\Delta$21',22' in 1.

Figure 4. An MM2 energy-minimized model of 1 displaying NOEs between H$_{-20}$/20' and H-21/21.

Compound 1 displayed optical activity with a similar sign and magnitude ([$\alpha$]$_{25}^{D} = -59^\circ$) to that of reported psammaplysins [2,4,6,8,9,11,13]. Therefore, it is more likely that psammaceratin A possesses the same biosynthetic path and shares similar stereochemistry at C-6/6' and C-7/7' with reported psammaplysins [2,4,6,8,9,11,13]. In addition, the sign and magnitude of the optical rotation of psammaceratin A are closely correlated to reported values [2,4,6,8,9,11,13]. The absolute configurations at C-6 and C-7 in psammaplysin A (2) were recently verified as 6R and 7R [23]. Therefore, we anticipate that psammaceratin A (1) shares the same absolute configurations of 6R,7R and 6'R,7'R with the parent compound, psammaplysin A [23]. Thus, psammaceratin A was assigned as (2Z,3Z)-2,3-bis(((3,5-dibromo-4-(3-((4R,5R)-8,10-dibromo-4-hydroxy-9-methoxy-
1,6-dioxa-2-azaspiro[4.6]undeca-2,7,9-triene-3-carboxamido)propoxy)phenethyl)amino)methylene)succinamide.

In an MTT assay [13,24], psammaceratin A (1) displayed the highest activity against HCT 116 cells with an IC_{50} value of 3.1 µM. On the contrary, psammaplysin A (2), with its free terminal amine moiety, was less active towards HCT 116, with an IC_{50} value of 5.1 µM. On the other hand, compound 2 was more active against MDA-MB-231 (IC_{50} = 3.90 µM), while 1 was less active against this cell line (IC_{50} = 5.25 µM). These data suggest that MDA-MB-231 and HCT 116 have high sensitivities towards 1 and 2, respectively. Finally, psammaceratin A (1) and psammaplysin A (2) displayed close and similar activity towards HeLa cells (IC_{50} = 8.50–9.40 µM) (Table 3) suggesting lower sensitivity of this cell line towards 1 and 2. Thus, psammaceratin A and psammaplysin A are considered as potential leads for the establishment of novel anticancer entities.

Table 3. Cytotoxic activities of 1 and 2.

| Compound | IC_{50} (µM) | MDA-MB-231 | HeLa | HCT 116 |
|----------|--------------|-------------|------|---------|
| 1        | 3.90 ± 0.20  | 8.50 ± 0.81 | 5.10 ± 0.41 |
| 2        | 5.25 ± 0.48  | 9.40 ± 0.89 | 3.10 ± 0.28 |
| 5-FU     | 13.0 ± 0.30  | 12.3 ± 0.25 | 4.60 ± 0.23 |

2.2. Structure of Psammaplysin A (2)

Psammaplysin A (2) (Figure 1) was identified by interpretation of its NMR and MS data and by comparison of its NMR data with the literature [4].

3. Materials and Methods

3.1. General Experimental Procedures

Optical rotations were acquired on a digital DIP-370 polarimeter (JASCO). The UV spectra were measured on a Hitachi 300 spectrometer. NMR data were acquired on a Bruker Avance DRX 600 MHz spectrometer using CD_{3}OD as the solvent. Positive ion HRESIMS spectra were collected on a Thermo LTQ Orbitrap XL mass spectrometer. A SiO_{2} RP HPLC column (Merck, 70–230 mesh ASTM) and Sephadex LH 20 (0.25–0.1 mm, Pharmacia) were used for chromatography. An HPLC column (AR II Cosmosil, Waters, 250 × 10 mm, 5 µm) was used for purification of the compounds.

3.2. Biological Materials

The Red Sea Pseudoceratina arabica (Keller, 1883) was harvested by hand via scuba down to −17 m from Anas Reef, Obhur at the Saudi Red Sea coast (N 021°39′17.5″, E 038°52′26.3″). The sponge consists of an encrusting mass of 1–2 cm with a conulose surface of yellowish green color underwater and greenish yellow interior. The sponge starts to turn blackish green in color after exposure to air. After storage in 70% ethanol solution, it turns completely into a black mass. The conules on the surface of the sponge are bluntly rounded, about 2–5 mm apart, and are of rubbery and compressible consistency. The specimen fragment measured about 12.0 × 5.0 × 1.5 cm. The sponge’s skeleton contains spare unequal fibers containing only pith. The outline branching is irregular, and the thickness measures 80–300 µm. The specimen corresponded to an Eritrean Red Sea specimen of P. arabica. The voucher was stored in the Zoological Museum’s collection at Amsterdam University under the code RMNHPOR 9161. Another specimen was stored at King Abdulaziz University under code #KSA-58.

3.3. Purification of 1 and 2

Freeze-dried sponge materials (230 g) were extracted thrice with 1000 mL MeOH. The combined extracts were dried under vacuum to afford 5.29 g of viscous residue. The extract was dissolved in MeOH–H_{2}O (6:4) and successively extracted with hexane, CH_{2}Cl_{2}, and...
EtOAc. The cytotoxic CH$_2$Cl$_2$ extract (2.1 g) was chromatographed over a SiO$_2$ VLC column using hexane/EtOAc/MeOH gradients to afford five fractions (Fr-1–Fr-5). Fractionation of the cytotoxic fraction (Fr-3) (270 mg) on a Sephadex LH 20 column with MeOH gave four major subfractions (Fr-3A–Fr-3C). The cytotoxic fraction Fr-3B (45 mg) was purified on an ODS HPLC column using 80% MeOH to yield compounds 1 (5.3 mg) and 2 (2.7 mg).

3.4. Spectral Data of the Compounds

Psammaceratin A (1): pale yellow solid; [α]$_D^{25}$ = −59° (c 0.1, MeOH); NMR data: Table 1; HRESIMS m/z 1624.6732, (calcd for C$_{48}$H$_{50}$Br$_4$N$_8$O$_{14}$Na, [M + Na]$^+$, 1624.6729).

Psammaplysin A (2) was identified by analyses of its 1D and 2D NMR data and by comparison of its spectroscopic data to those in the literature [4].

3.5. Cytotoxicity of the Compounds

3.5.1. Preparation of Cell Lines and Cell Culture

The human cell lines HCT116 (colorectal carcinoma, ATCC CCL-247), MDA-MB-231 (triple-negative breast cancer, ATCC HTB-26), and HeLa (human cervical carcinoma, ATCC CCL-2) were used in this investigation. For MDA-MB-231, DMEM with 1% penicillin–streptomycin and 10% FBS was used, while RPMI 1640 medium with 10% FBS and 1% penicillin–streptomycin was used for culturing HCT116 and HeLa cells. All cells were incubated at 37 °C with 5% CO$_2$ and 95% humidity.

3.5.2. Cytotoxicity and Antiproliferative Activity

The evaluation of the cytotoxicity of the compounds was carried out via MTT assay as previously described [13,24]. Briefly, cells were incubated at 37 °C overnight in 5% CO$_2$/air, followed by the addition of the compounds at the top row of a 96-well microtiter plate and descending serial dilution (1:4) of the concentration. The cells were incubated for 72 h with the compounds. Subsequently, the cell viability was estimated at 490 nm on a Molecular Devices Emax microplate reader using the Cell Titer 96 AQueous non-radioactive cell proliferation protocol. The IC$_{50}$ values of the compounds (expressed in micromoles) were evaluated using the program SOFTmax PRO. 5-Fluorouracil (5-FU) and DMSO were used as positive negative controls.

4. Conclusions

Bioassay-directed fractionation of the cytotoxic extract of the Red Sea sponge *Pseudoceratina arabica* afforded an unprecedented psammaplysin dimer, psammaceratin A (1), along with psammaplysin A (2). Psammaceratin A, with its unique (2Z,3Z)-2,3-bis(aminomethylene)succinamide backbone connecting two units of psammaplysin A, represents the first dimer of this type within the psammaplysin family. This previously unknown functional group, 2,3-bis(aminomethylene)succinamide, offers a novel synthetic moiety that could be utilized as an isostere in synthetic chemistry, as a novel connecting moiety, or for other design and derivatization purposes. Accordingly, psammaceratin A and psammaplysin A are potential scaffolds for the development of novel antitumor leads.

**Supplementary Materials:** The following are available online at https://www.mdpi.com/article/10.3390/md19080433/s1, Figures S1–S9: HRESIMS, $^1$HNMR, $^{13}$C NMR, DEPT, COSY, HSQC, and HMBC, and NOESY spectra of psammmaceratin A (1).

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