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The total synthesis and functional evaluation of fourteen stereoisomers of yaku'amide B. The importance of stereochemistry for hydrophobicity and cytotoxicity[†]

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Yaku'amide B is a highly unsaturated linear tridecapeptide and an extremely potent cytotoxin. Herein, we describe the synthesis of fourteen new stereoisomers of yaku'amide B using a unified assembly strategy. The hydrophobicities and cytotoxicities of these analogues were analyzed, along with those of four previously prepared isomers. Although all of the analogues share a common planar structure, their log *D* values varied significantly (3.39–5.32), presumably reflecting their distinct three-dimensional shapes. Subnanomolar-level cytotoxicity was observed for the natural yaku'amide B and its epimer of the N-terminal acyl group, whereas the other sixteen isomers exhibited 13- to 1200-fold weaker activities than that of the natural isomer. These data indicated the importance of the overall stereostructure of the 13-mer sequence of yaku'amide B for exerting its potent toxicity.

Introduction

Yaku'amides A (1) and B (2) were isolated from a deep sea sponge *Ceratopsion* sp. (Fig. 1) and were reported to exhibit exceptionally potent cytotoxicity towards P388 murine leukemia cells.¹ Cytotoxicity assays of 1 using a panel of 39 human cancer cell lines (JFCR39)² unveiled its distinct growth-inhibitory profile in comparison with 38 clinically available anticancer drugs, suggesting its potentially novel mode of action. The linear tridecapeptide sequences of 1 and 2 only differ in their third residue (Gly-3 for 1 and Ala-3 for 2), and both compounds contain four tetrasubstituted α , β -dehydroamino acids³ (Δ Ile-2, Δ Ile-4, Δ Ile-9, and Δ Val-13) and seven other non-proteinogenic amino acids. Additionally, an N-terminal acyl (NTA) group and a C-terminal amine (CTA) cap the N- and C-termini,

respectively. These unusual structural features and potent biological activities of **1** and **2** have spurred interest in the chemical and biological communities.⁴

Since only minute amounts of 1 and 2 have been obtained from natural sources (1: 1.3 mg; 2: 0.3 mg), their structural determination is incomplete and detailed biological studies have been hampered. To address these issues, we launched synthetic studies of 1 and 2 as the initial phase for further investigations of their structures and bioactivities. Consequently, the C4-epimers of the NTA moiety of the proposed structures 1a/b and 2a/b were chemically constructed by developing a new protocol for construction of the α,β -unsaturated amino acid residues.^{5,6} Namely, Cu-catalyzed C(sp²)-N bond formation between the primary amides and the alkenyl iodides was adopted to construct four building blocks with the tetrasubstituted enamide moieties (4a, 5a, 9, and 10a).⁷ Then, the enamide building blocks and the protected amino acids were condensed stepwise to furnish the possible proposed structures 1a/b and 2a/b. However, both the NTA-epimers 1a/b and 2a/b exhibited different chromatographic behaviors from naturally occurring 1 and 2, respectively. This finding prompted investigations directed toward determining the true structures of 1 and 2. Structural information was gathered through degradative studies of 0.05 mg of the available 2 and synthetic preparation of all the possible structures of the degraded partial structures. These studies allowed us to establish the structures of the natural 1 and 2 to be 1c and 2c. Both the revised structures possess the inverse configurations at the C_{α} positions of the four amino acid residues (OHVal-7, -8, Val-11, and -12) of the proposed structures, and the S-configuration at C4 of NTA. Compounds 1c and 2c were then prepared by total syntheses to confirm their structural authenticity.

During the process of structural revision by a combination of degradation and synthesis, the total syntheses of the multiple potential isomers of the natural **2** were independently performed without achieving the revised structure **2c**. Specifically, the fourteen analogues **2e-r** having different amino acid stereochemistries from those of the proposed structures **2a/b**

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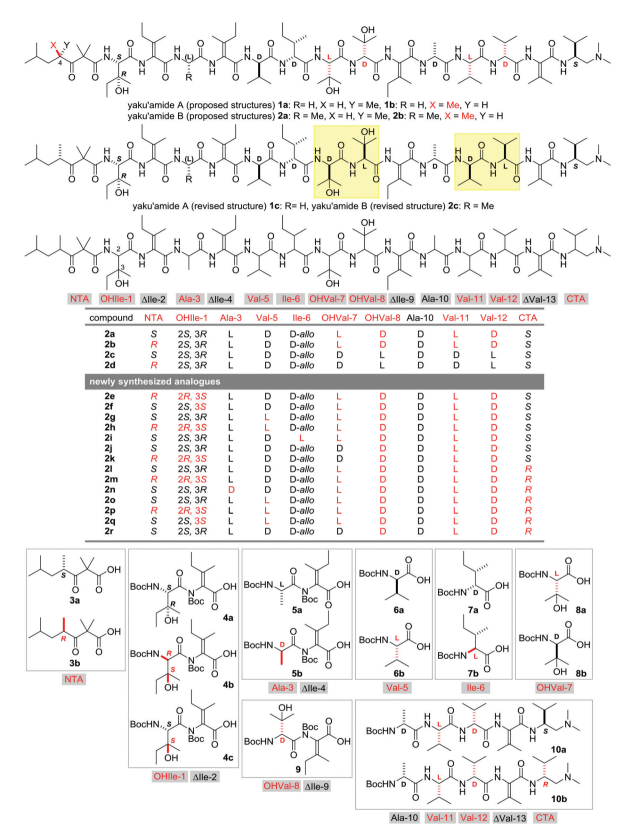


Fig. 1 Structures of 1a/b (yaku'amide A proposed structures), 1c (yaku'amide A revised structure), 2a/b (yaku'amide B proposed structures), 2c (yaku'amide B revised structure), 2d (NTA-epimer of yaku'amide B), newly synthesized analogues 2e-r, and the synthetic fragments 3-10. The bonds displayed in red indicate the altered stereochemistry from the natural 2c. Boc = tert-butoxycarbonyl; $\Delta IIe = \alpha, \beta$ -dehydroisoleucine; $\Delta VaI = \alpha, \beta$ -dehydrovaline; OHIV = β -hydroxysoleucine; OHV = β -hydroxyvaline.

were synthesized. Upon analysis by ultrahigh-performance liquid chromatography (UHPLC), all of the analogues 2e-r provided retention times distinct from that of the natural 2, indicating that they did not correspond to the true structure. Although the syntheses of 2e-r did not contribute to the structural elucidation of 2c, this focused library of stereoisomers provided us with a valuable opportunity for investigating the structure-activity relationship (SAR) of yaku'amide B (2c).⁸ Compounds 2e-r share the same planar structure and thus their functional evaluations would clarify the stereochemical requirement of the amino-acid residues for the unique physicochemical and potent biological properties of 2c. Furthermore, the SAR study would offer useful insight into the unknown mode of action of 2c. Herein, we report the syntheses and functional analyses of the fourteen analogues 2e-r of yaku'amide B (2c). These studies showed that the configurational changes of the main chain of 2c significantly affected the hydrophobicities and greatly diminished the cytotoxicities. Only 2c and its C4-epimer of NTA 2d displayed subnanomolarlevel cytotoxicity towards P388 cells, demonstrating the biological importance of the main chain stereochemistry.

Results and discussion

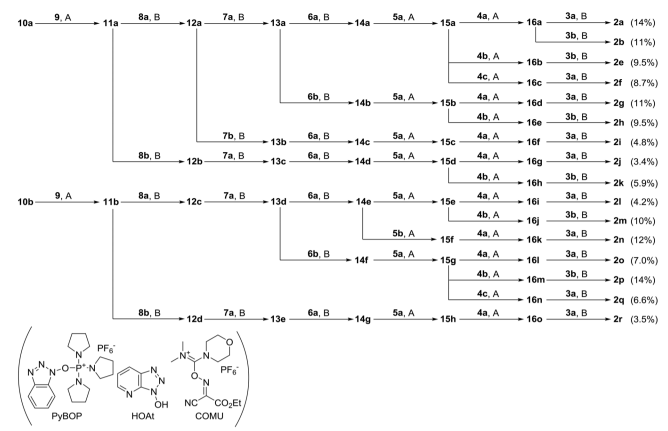
The fourteen new analogues of yaku'amide B, 2e-r, have different configurations from the natural 2c at NTA, OHIle-1, Ala-3, Val-5, Ile-6, OHVal-7, OHVal-8, Val-11, Val-12, and/or CTA (Fig. 1), while the double-bond geometries of Δ Ile-2, -4, and -9 and the D-stereochemistry of Ala-10 are retained. As 2e-r were designed based on the proposed structures 2a/b, all of these peptides possess the same stereochemistries at OHVal-8, Val-11, and Val-12. We previously reported the total syntheses of 1a-c and 2a-d by assembling eight building blocks. This efficient strategy was applied to the syntheses of 2e-r by switching the fragments to their corresponding stereoisomers. Accordingly, sixteen fragments were necessary for the construction of 2e-r (NTA: 3a and 3b; OHIle-1- Δ Ile-2: 4a, 4b, and 4c; Ala-3-∆Ile-4: 5a and 5b; Val-5: 6a and 6b; Ile-6: 7a and 7b; OHVal-7: 8a and 8b; OHVal-8-ΔIle-9: 9; Ala-10-Val-11-Val-12-∆Val-13-CTA: 10a and 10b). While 3a/b, 4a, 5a, 6a/b, 7a/b, 8a/b, 9 and 10a/b were previously prepared on the way toward the total synthesis and structural elucidation of 1c and $2c_{5}^{5}$ dipeptides 4b (enantiomer of 4a), 4c (diastereomer of 4a) and 5b (enantiomer of 5a) were newly obtained by using Cu-mediated C(sp²)–N bond formation from the corresponding monomers.⁹

To produce the analogues 2e-r, the sixteen synthetic fragments were condensed according to the assembly procedure that was optimized for the route to 2a and 2b^{5*a*} (Scheme 1). The unified total syntheses of the known 2a/b and the new 2e-rstarted with the TFA-promoted removal of the N_{α} -Boc group from 4-mer derivatives **10a** and **10b**. The obtained two amines were separately treated with carboxylic acid **9** in the presence of PyBOP ¹⁰/HOAt ¹¹ to generate the protected 6-mer peptides **11a** and **11b**, respectively. Detachment of the two Boc groups of OHVal-8 and Δ Ile-9 of **11a** and **11b** with TFA afforded the

corresponding amines, which were amidated with 8a or 8b by the action of COMU,¹² giving rise to the four 7-mer derivatives 12a/b and 12c/d, respectively. Stepwise elongation from 12a-d using the five building blocks completed the syntheses of 2a/ b/e-r. Namely, two cycles of TFA-mediated N_{α} -deprotection and COMU-promoted condensation using 7a/b and 6a/b converted 12a-d to the seven 9-mer peptides $(12a-d \rightarrow 13a-e \rightarrow 13a-e)$ 14a-g). The obtained compounds 14a-g were then transformed to fifteen 13-mer peptides 16a-o through repeated Boc removal and PyBOP/HOAt-promoted conjugation using 5a/b or 4a-c (14a-g \rightarrow 15a-h \rightarrow 16a-o). Finally, COMU enabled the coupling of the NTA unit 3a/b with 16a-o, delivering the sixteen yaku'amide analogues 2a, 2b, and 2e-r. As a result of the present divergent synthesis, the fourteen new peptides 2e-r were efficiently constructed in 14 steps from 9 in reasonable yields (2a, 2b, 2e, 2f, 2g, 2h, 2i, 2j, 2k, 2l, 2m, 2n, 2o, 2p, 2q, and 2r for 14, 11, 9.5, 8.7, 11, 9.5, 4.8, 3.4, 5.9, 4.2, 10, 12, 7.0, 14, 6.6, and 3.5%, respectively). These achievements corroborated the high adaptability of our synthetic strategy for the preparation of diverse yaku'amide isomers.

The completion of the total syntheses of 2a-d and the fourteen new analogues 2e-r permitted us to conduct systematic functional analyses. Since the overall hydrophobicity influences the molecular behavior and interaction with the cell membrane and biomolecules, it is one of the most critical factors affecting the biological activity of a compound. Accordingly, the hydrophobicities of isomers 2a-r were estimated by determination of their octanol/water distribution coefficients $(\log D)$.¹³ To calculate the $\log D$ values of 2a-r, the retention times of 2a-r and the standard samples with known $\log D$ values were compared under identical UHPLC conditions employing an ODS column.¹⁴⁻¹⁶ Although all the peptides possess the same planar structure, the numbers of log D varied significantly, from 3.39 to 5.32 (Table 1), where the values of the natural 2c (4.13) and NTA-epimer 2d (4.34) were close to the average. As the amino acid monomers at the same residue numbers of 2a-r are enantiomeric with the exception of several diastereomeric monomers (OHIle-1 of 2f/q and Ile-6 of 2i), the hydrophobicities of most individual monomers are identical. Therefore, the diverse log D values of 2a-r would originate from differences in the entire three-dimensional structure of each compound.¹⁷ Specifically, the main-chain stereochemistries of 2a-r would organize the locations and orientations of the hydrophobic side-chains of the molecules, thereby changing the hydrophobic interactions between the peptides and the ODS column. Thus, quantification of the log D values demonstrated the significant effect of configurational alterations on the hydrophobicity and the molecular shape of this family of compounds.

Next, the cytotoxicities of the synthetic analogues against P388 mouse leukemia cells were assessed using the XTT assay method (Table 1).¹⁸ The natural isomer **2c** exhibited subnanomolar-level cytotoxicity, with an IC₅₀ value of 0.51 nM. Whereas the stereochemical inversion of the C4 of NTA (**2d**) induced no drastic change in cytotoxicity (IC₅₀ = 0.95 nM), alteration of the main chain stereochemistry decreased the



Scheme 1 Syntheses of the analogues 2a, 2b, and 2e–r. Overall yields from 9 are given in parentheses. Reagents and conditions: (A) (i) TFA, CH_2Cl_2 , (ii) PyBOP, HOAt, *i*-Pr₂NEt, DMF; (B) (i) TFA, CH_2Cl_2 , (ii) COMU, 2,4,6-collidine, DMF. COMU = (1-cyano-2-ethoxy-2-oxoethylidenaminooxy)dimethyl-amino-morpholino-carbenium hexafluorophosphate; DMF = N,N-dimethylformamide; HOAt = 1-hydroxy-7-azabenzotriazole; PyBOP = benzotri-azol-1-yl-oxytripyrrolidinophosphonium hexafluorophosphate; TFA = trifluoroacetic acid.

 $\label{eq:table_$

Compound	$\log D^a$	$\operatorname{IC}_{50}^{b}(\mathrm{nM})$
2a	3.48	98
2b	3.39	410
2c	4.13	0.51
2d	4.34	0.95
2e	3.57	620
2f	3.56	450
2g	3.99	92
2h	3.91	530
2i	3.39	300
2j	4.25	9.3
2k	4.40	34
21	3.93	23
2m	4.11	93
2n	4.09	73
20	3.92	45
2p	3.90	260
2q	4.06	100
2r	5.32	6.8

 $^{a} \log D$ values were measured by UHPLC methods. b IC₅₀ values were determined from the results of growth inhibition assays (XTT method) performed on P388 mouse leukemia cells. XTT = 3'-[1-[(phenylamino)-carbonyl]-3,4-tetrazolium]bis(4-methoxy-6-nitro)-benzenesulfonic acid hydrate.

cytotoxicities of the peptides by 13- (6.8 nM, 2r) to 1200-fold (620 nM, 2e). Interestingly, 2j and 2r that have the identical left half structure (NTA to OHVal-7) to the natural 2c are the only analogues among 2a/b/e-r with nanomolar-level toxicities. On the other hand, the toxicities of the originally proposed structures 2a and 2b were 190 and 800 times weaker than those of 2c, supporting the validity of the revised structure. Overall, the greater potency of 2c/d compared to the other fourteen isomers indicated the biological importance of the configurations of the thirteen amino-acid residues of 2c/d.

To investigate the relationship between the physicochemical and biological properties, the log *D* values of the eighteen yaku'amide isomers **2a–r** were plotted against their IC₅₀ numbers (Fig. 2). Consequently, these two values of the sixteen non-natural analogues **2a**, **2b**, and **2e–r** were found to have a modest linear relationship with the more hydrophobic analogues among **2a**, **2b** and **2e–r** generally exhibiting a higher cytotoxicity. Because of the variable stereostructures of **2a**, **2b** and **2e–r**, the cytotoxicities of these non-natural analogues would relate to their non-specific hydrophobic interactions with membranes or biomolecules. In contrast, the naturally occurring isomer **2c** and its NTA epimer **2d** located well beyond the approximate straight line of the other sixteen ana-

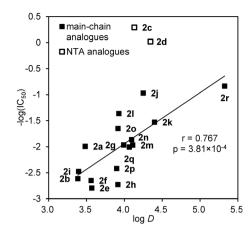


Fig. 2 Plot of $-\log(|C_{50})$ values against $\log D$ values of 2a-r. The black line represents the linear fit of the main-chain analogues 2a, 2b and 2e-r. Correlation coefficient (r) and p-value are also displayed.

logues. For instance, the cytotoxicities of 2c and 2d were significantly higher than those of 2j and 2k, despite their similar log *D* values. Taken together, these data suggested a mode of action of 2c and 2d distinct from that of 2a, 2b and 2e-r. Hence, it is likely that peptides 2c and 2d display their bioactivities through specific binding to a chiral biomolecule rather than through non-specific hydrophobic interactions. According to this hypothesis, the main chain stereochemistry of 2c and 2d would fold into the appropriate three dimensional structure, form a complementary surface to a target biomolecule, and thereby cause a potent toxic effect on cancer cells.

Conclusions

The fourteen analogues 2e-r of the naturally occurring yaku'amide B (2c) were newly synthesized in a divergent fashion by assembling stereoisomers of the eight fragments 3-10. The physicochemical and biological properties of the prepared 2e-r were systematically evaluated and analyzed along with the originally proposed structures 2a/b, the natural 2c, and the C4epimer of NTA 2d. Although all the isomers 2a-r share the same planar structure, they showed diverse hydrophobicities and cytotoxicities against P388 mouse leukemia cells. The natural 2c and the NTA-epimer 2d had average hydrophobicity values $(\log D)$, yet they were 13–1200 times more cytotoxic compared to the sixteen non-natural isomers 2a, 2b, and 2e-r. The linear correlation between the $\log D$ and IC_{50} values of the weaker analogues 2a/b/e-r suggested that the bioactivities of these analogues would be affected through non-specific hydrophobic interactions. On the other hand, the excellent potency of 2c and 2d would originate from a mode of action distinct from that of the other isomers. Thus, the main chain stereochemistries of 2c and 2d would be important for folding into a specific threedimensional structure, which binds to a target biomolecule of the cell to display its cytotoxicity. Detailed biological studies to prove this hypothesis are currently underway in our laboratory.

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