PHE¹² of deacetyl-thymosin β_4 is one of the structural essentials for restorative effect on the impaired blastogenic response of uraemic T-lymphocytes. In order to evaluate the functional roles of this phenyl group in the restorative effect on impaired T-lymphocytes, two analogues, [1-Nal¹²]deacetyl-thymosin β_4 and [Cha¹²]deacetyl-thymosin β_4 , were synthesized by a solid-phase method and evaluated for restorative effect on the impaired blastogenic response of uraemic T-lymphocytes. The results indicated that [1-Nal¹²]deacetyl-thymosin β_4 which had a bulky naphthyl ring showed a stronger restorative effect than that of deacetyl-thymosin β_4 , but it was slightly weaker than that of [Phe(4F)¹²]deacetyl-thymosin β_4 . However, [Cha¹²]deacetyl-thymosin β_4 showed no restorative effect on the impaired blastogenic response of uraemic T-lymphocytes.

Key words: Blastogenic response, Deacetyl-thymosin β_4 analogue synthesis, Impaired Tlymphocyte, Restorative effect, Uraemic patient

Functional roles of Phe¹² of deacetyl-thymosin β_4 in the impaired blastogenic response of uraemic T-lymphocytes

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Abbreviations

Boc, tert-butyloxycarbonyl; tBu, tert-butyl; DMF, N,Ndimethylformamide; TFA, trifluoroacetic acid; Fmoc, 9-fluorenylmethoxycarbonyl; HOBT, 1-hydroxybenzotriazole; DCC, dicyclohexylcarbodiimide; HBF4, tetrafluoroboric acid; EDT, ethane-1,2-dithiol; AcOH, acetic acid; HPLC, high-performance liquid chromatography; TLC, thin-layer chromatography; 1-Nal, 1-naphthylalanine; Cha, cyclohexylalanine; PHA, phytohaemagglutinin; RPMI, Rosewell Park Memorial Institute; SDS, sodium dodecyl sulphate; PBS, phosphate-buffered saline; FCS, fetal calf serum; FAB-MS, fast atom bombardment mass spectrometry; Pam, phenylacetoamido-methyl; Ac, acetyl.

Introduction

The impairment of immunological responsiveness in uraemic patients is well known. All aspects of the immune response appear to be affected by the uraemic state. The numbers, subpopulations and reactivities of circulating lymphocytes may be altered by uraemia. This impairment has been implicated in easy susceptibility to infections and increased incidence of malignancy.

Thymosin β_4 , an N-terminal acetylated peptide containing 43 amino acid residues, was first

isolated from calf thymus by Low *et al.*³ and has the following amino acid sequence: Ac-Ser-Asp-Lys-Pro-Asp-Met-Ala-Glu-Ile-Glu-Lys-Phe-Asp-Lys-Ser-Lys-Lys-Thr-Glu-Thr-Glu-Glu-Lys-Asn-Pro-Leu-Pro-Ser-Lys-Leu-Lys-Glu-Thr-Ile-Glu-Glu-Glu-Lys-Gln-Ala-Gly-Glu-Ser-OH. This peptide exhibits several biological activities that are important for maturation and functioning of the immune systems.⁴

Previously⁵⁻⁷ we reported syntheses of deacetyl-thymosin β_4 and its fragments and that some of the fragments could have a restorative effect on the impaired cell-mediated immunological functions. We also noticed that the acetyl group at the N-terminal serine residue of thymosin β_4 , is not required for the restorative effect on the impaired cell-mediated immunological functions.⁵

In an earlier paper,⁸ we reported that the synthetic [Phe(4F)¹²]deacetyl-thymosin β_4 exhibited stronger restorative effect on the impaired blastogenic response of Tlymphocytes isolated from uraemic patients than that of our synthetic deacetyl-thymosin β_4 . In this study, the strong electron-withdrawing fluoride atom on the para-position of the aromatic ring results in an analogue that possesses stronger activity than that of the parent molecule.⁸ This result seems to suggest that modification of the Phe residue of thymosin β_4 could produce more potent analogues capable of a restorative effect

on impaired blastogenic response of Tlymphocytes.

The purpose of the present study was to synthesize two thymosin β_4 analogues, [1-Nal¹²]deacetyl-thymosin β_4 and [Cha¹²]deacetyl-thymosin β_4 by the solid-phase method and to compare the restorative effect of these two analogues on the impaired blastogenic response of uraemic Tlymphocytes.

Materials and Methods

Fmoc-amino acid derivatives and Fmoc-Ser (tBu)-Pam-resin (0.64 mmol/g, 100-200 mesh) were purchased from Kokusan Chemical Works Itd (Japan), Watanabe Chemical Industries Itd (Japan), Peptide Institute Inc (Japan) and Sigma Chemical Co. (USA). TIC was effected with silica gel (Kieselgel $60F_{254}$, Merck) on precoated aluminium sheets using n-BuOH-AcOH-pyridine- H_2) = 4:1:1:2 as a solvent system. Analytical HPIC and amino acid analysis were conducted with a Shimadzu IC6A and Hitachi 835A, respectively. The FAB-MS spectrum was obtained on a VG analytical 2AB-2SEQ spectrometer equipped with the 11-250J data system.

Solid-phase peptide synthesis

Peptide synthesis was performed manually by the stepwise solid-phase method with a handmade peptide synthesizer, using the base-labile Fmoc group for protecting the \alpha-amino groups, and such acid-labile groups as the tBu for the hydroxy and carboxy groups, the Boc for the εamino groups of Lys, and the sulphoxide for Met. The peptide was assembled on Fmoc-Ser (tBu)-Pam-resin. The Emoc group was removed with 30% piperidine in DMF. Elongation of the peptide chain was carried out by the DCC-HOBT method in CH₂Cl₂-DMF (1:1) or in Nmethyl-2-pyrrolidone. The coupling reaction and deprotection of the Fmoc group were monitored by the ninhydrin test. The general procedure for each synthetic cycle (as a starting mateial 0.64 mmol/g of Fmoc-Ser(tBu)-Pamresin: 400 mg) was: (1) CH₂Cl₂ wash (twice); (2) DMF wash (twice); (3) deprotect: DMFpiperidine (7:3) for 20 min; (4) DMF wash (twice); (5) dioxane-water (2:1) wash (twice); (6) DMF wash (three times); (7) CH₂Cl₂ wash (three times); (8) addition of 3 eq Fmoc-amino acid, HOBT, and DCC in CH₂Cl₂-DMF (1:1) or N-methyl-2-pyrrolidone; (9) add 1.0 ml of diisopropylethylamine in CH₂Cl₂; (10) reaction for 120 min; (11) recoupled if necessary by repeating steps 7-10; (12) DMF wash (three times); (13) isopropanol wash (three times); (14)

CH₂Cl₂ wash (four times). Whenever the ninhydrin test was still slightly positive, even after three couplings, the remaining unreacted amino groups were acetylated with Ac₂O-pyridine in DMF. The peptide resin (200 mg) was treated with 2 M HBF₄-thioanisole in TFA (7 ml) in the presence of m-cresol (218 µl, 100 eq) and EDT (524 µl, 300 eq) at 4°C for 90 min.

After the deprotection, the resin was removed by filtration and the filtrate was evaporated under reduced pressure and the residue was solidified by addition of anhydrous peroxide free ether to give a crude peptide. The resulting powder was dissolved in H₂O (6 ml). The solution was treated with Amberlite CG4B (acetate form, approximately 3 g) for 30 min, and filtered by suction and evaporated in vacuo. The residue was dissolved in H2O (10 ml). The solution, after addition of dithiothreitol (20 mg), was incubated at 60°C under N_0 gas for 36 h. The solvent was evaporated off *in vacuo* and the residue was dissolved in a small amount of 1%AcOH and then applied to a column of Sephadex G-25 (2.3 \times 95 cm), which was eluted with the same solvent. Individual fractions (5 ml each) were collected and absorbancy at 230 nm was determined for each fraction. The fractions corresponding to the front main peak were combined and the solvent was removed by lyophilization. The peptide was further purified by semi-preparative PR-HPLC. The semi-preparative PR-HPLC was performed on a Nucleosil C18 column $(250 \times 10 \text{ mm I.D.}; 7 \mu\text{m particle size}; \text{Macherey})$ Nagel). Solvent A was 0.05% TFA in water and solvent B was 60% acetonitrile in solvent A. A linear gradient was applied from 10 to 50% B during 50 min, at a flow rate of 3.0 ml/min. Detection of the peptide was set at 230 nm. The major peak was lyophilized to give the purified product. [1-Nal¹²] deacetyl-thymosin β₄; 20.3 mg (20% calculated from the starting Cterminal amino acid). [Cha¹²]deacetyl-thymosin β_4 ; 22.6 mg (23%) calculated from the starting Cterminal amino acid) (Fig. 1).

Patient selection

Three uraemic patients who needed dialysis treatment three times a week and were suffering from recurrent infectious diseases (pneumonia and tuberculosis) were selected. Examination of cellular immunocompetence of these patients revealed a significant decrease in blast-formation by PHA. ³H-thymidine incorporation values of these patients were 11826, 12042 and 12153 cpm respectively (normal values: 41195–42659 cpm).

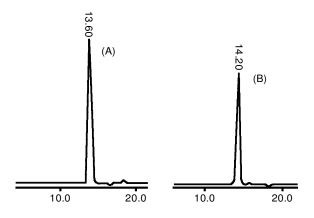


Fig. 1. HPLC profiles of (A) [1-Nal 12]deacetyl-thymosin β_4 and (B) [Cha 12]deacetyl-thymosin β_4 .

Venous blood was obtained from these uraemic patients for the fluorometric blast-formation test. Venous blood samples from three healthy donors were used as a control. The fluorescence excitation spectrum was measured with an Oyo-Bunko ULOG-FLOUSPEC 11A fluorometer. Kits for the fluorometric blast-formation test were purchased from Japan Immunoresearch Laboratories Co. Ltd (Japan).

Fluorometric blast-formation test

A 3 ml aliquot of venous blood was drawn into a syringe containing 25 U/ml of heparin and then mixed with 3 ml of PBS. Lymphocytes were isolated in a Hypaque-Ficoll gradient. lymphocytes Isolated were adjusted 1.0×10^6 /ml with PBS. The lymphocytes were cultured in 0.5 ml of RPMI 1640 (Gibco) with 10% FCS (Dainippon Pharmaceutical Co.) in microplates. Cultures of each combination were incubated at 37°C in the presence of one of the peptides in a humidified atmosphere of 5% CO₂ in air or 12 h and then PHA (0.125%) was added to each well and incubation was continued under the same conditions for 60 h. Lymphocytes in each well were transferred into a test tube and centrifuged for 10 min at 240 g, then the supernatant was removed. A 2 ml aliquot of 0.125% SDS was added to the residue and stirred for 20 min at room temperature; lymphocytes were completely destroyed and solubilized by this procedure. Ethidium bromide solution

(2 ml) was added to the above solution and the mixture was stirred for 15 min at room temperature. The fluorescence excitation spectrum was measured as previously described.⁷

Results

In order to construct the peptide chain, the Fmoc-based solid-phase method was employed. Fmoc-Ser(tBu)-Pam-resin was placed in the reaction vessel and the combination of piperidine treatment and DCC plus HOBT procedure served to elongate the peptide chain manually according to the usual method.

We encountered no serious difficulties during the elongation of the entire sequences of the two analogues, although the double coupling procedure was employed when the resin became positive to the ninhydrin test, after a single coupling. The coupling cycle included a capping step with acetic anhydride (5 min) to prevent the formation of deleted sequences. The amino acid compositions of the protected peptide resins thus assembled were in good agreement with those predicted by theory after acid hydrolysis with 12 N HQ-proprionic acid (1:1). The protected peptide resins thus obtained were then treated with 2 M HBF₄-thioanisole in TFA at 4°C for 90 min to cleave the peptide chain from the resin and at the same time to remove all side-chain protecting groups employed. The Met(O) residue was reduced back to Met in two steps, firstly with 2 M HBF $_4$ thioanisole in TFA during the above acid treatment, and secondly with dithiothreitol during incubation of the unprotected peptide.

The crude peptides thus obtained were then successively purified by gel-filtration on Sephadex G-25 and semi-preparative HPLC. The two purified peptides exhibited single peaks on analytical HPLC. The two products possessed amino acids in ratios consistent with those predicted from the sequences of the two analogues after acid hydrolysis. The homogeneity of the peptides was checked by TLC, HPLC, amino acid analysis after 6 N HCl hydrolysis, and FAB-MS spectrometry. Physicochemical data for the synthetic analogues are shown in Tables 1 and 2.

Table 1. Characterization of synthetic thymosin β_4 analogues

Peptide	Yield ^a (%)	$[\alpha]_D^{21}$ (c = 0.5, 1% AcOH)	TLC ^b Rf	FAB-MS ^c (MH ⁺)
[1-Nal ¹²]deacetyl-thymosin β_4 [Cha ¹²]deacetyl-thymosin β_4	20	−84.9°	0.09	4971.13
	23	−79.5°	0.11	4927.32

^aFinal yield after deblocking and purification starting from Fmoc-Ser(tBu)-Pam-resin.

^bSee the experimental section.

[°]Found values were in agreement with calculated values.

Table 2. Amino acid analysis of synthetic thymosin β_4 analogues^a

Peptide	Gly	Ala	Leu	lle	Pro	Ser	Thr	Met ^b	Lys	Asp	Glu	1-Nal	Cha
[1-Nal ¹²]deacetyl-thymosin β_4 [Cha ¹²]deacetyl-thymosin β_4	1.00 1.00	1.96 1.92						0.92 0.93			10.38 10.81	0.94	0.96

^aAfter acid hydrolysis with 6 N HCl at 126°C for 25 h.

Table 3. Effects of synthetic deacetyl-thymosin β_4 and its analogues on the impaired PHA stimulation of uraemic T-lymphocytes

Peptide	No. of determinations	Dose (μg/ml)	Sl ^{a,b}
c	3	_	274.2 ± 49.8
<u>d</u>	3		114.3 ± 48.7^{f}
Deacetyl-thymosin β ₄ d,e	3	0.1	115.1 ± 49.3
Deacetyl-thymosin β ₄ ^{d,e}	3	1.0	198.3 ± 50.1^{g}
[Phe(4F) ¹²]deacetyl-thymosin β ₄ d,e	3	0.1	190.6 ± 48.5^{g}
[Phe(4F) ¹²]deacetyl-thymosin $\beta_4^{d,e}$	3	1.0	230.5 ± 48.3^{g}
[1-Nal ¹²]deacetyl-thymosin β ₄ d,e	3	0.1	179.7 ± 49.6^{g}
[1-Nal ¹²]deacetyl-thymosin β ₄ d,e	3	1.0	216.8 ± 48.4^{g}
[Cha ¹²]deacetyl-thymosin β ₄ ^{d,e}	3	1.0	117.9 ± 49.0
[Cha ¹²]deacetyl-thymosin β ₄ d,e	3	10.0	115.3 ± 48.9

^aEach value represents the mean \pm SD of triplicate measurements.

The immunological effects of the synthetic deacetyl-thymosin β_4 , [Phe(4F)¹²]deacetyl-thymosin β_4 , and [Cha¹²]deacetyl-thymosin β_4 were examined by the JIMRO (Japan Immunoresearch Laboratories Itd) fluorometric blast-formation test. Responses of Tlymphocytes to mitogenic stimulation were significantly lower in uraemic patients that were those of normal persons. The *in vitro* effect of the synthetic peptides on the impaired PHA response of Tlymphocytes from uraemic patients is shown in Table 3.

Discussion

Comparison of the stimulation index (SI) values of the blastogenic transformation of Tlymphocytes into lymphoblasts with mitotic activity upon PHA stimulation shows that in the case of the uraemic patients investigated, the synthetic analogue, [1-Nal¹²]deacetyl-thymosin β_4 which had a bulky naphthyl ring exhibited stronger restorative activity than that of our synthetic deacetyl-thymosin β_4 , but it was a little bit weaker than that of [Phe(4F)¹²]deacetyl-thymosin β_4 . However, the synthetic [Cha¹²]deacetyl-

thymosin β_4 had no restorative effect even at a much higher concentration (Table 4).

Those results exhibited that not only 4-fluor-ophenyl ring of deacetyl-thymosin β_4 but also more bulky naphthyl ring of deacetyl-thymosin β_4 could bind with receptors of Tlymphocytes more strongly than a phenyl ring of deacetyl-thymosin β_4 . On the contrary, another analogue, [Cha¹²] deacetyl-thymosin β_4 which contains an aliphatic ring at position of 12 instead of an aromatic ring showed no restorative effect. This result seems to suggest that aromaticity at position of 12 of thymosin β_4 plays significant roles for restorative activity on impaired blastogenic response of Tlymphocytes.

Table 4. Relative potencies of synthetic deacetyl-thymosin β_4 and its analogues on the impaired PHA stimulation of T-lymphocytes of uraemic patients

Peptide	Relative potency (molar basis)
Deacetyl-thymosin β_4 [Phe(4F) ¹²]deacetyl-thymosin β_4 [1-Nal ¹²]deacetyl-thymosin β_4	1.00 10.48 9.86
[Cha ¹²]deacetyl-thymosin β ₄	

bMet + Met(O).

 $[^]bSI$ (stimulation index) was calculated according to the following formula: SI = $(l_2-l_0)/(l_1-l_0)\times 100$, where l_2 is mean fluorescence intensity of PHA-activated lymphocytes, l_1 is mean fluorescence intensity of PHA-nonactivated lymphocytes and l_0 is mean fluorescence intensity of ethidium bromide.

^cNormal venous lymphocytes.

dUraemic patients' lymphocytes.

elncubation was carried out at 37°C in a humidified atmosphere of 5% CO₂ in air for 12 h in the presence of each peptide.

 $^{^{\}dagger}P$ < 0.03, when compared with the normal subject using Student's t-test.

 $^{^{9}}P < 0.01$, when compared with the uraemic patients using Student's t-test.

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