

NEW METHOD FOR THE SYNTHESIS OF DEHYDROTRIPEPTIDES,
USING N-CARBOXY α -DEHYDROAMINO ACID ANHYDRIDE AS SYNTHON

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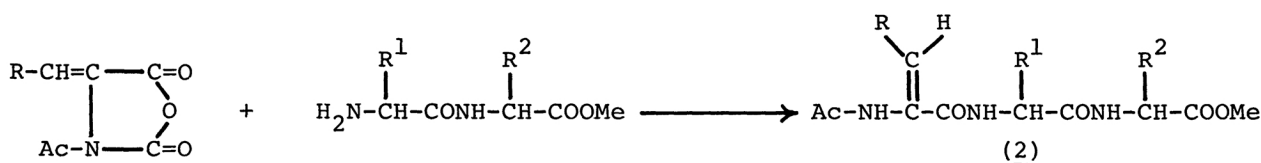
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Various dehydrotripeptides, containing one or two α -dehydro-amino acid residues, were readily synthesized by using N-carboxy α -dehydroamino acid anhydride as synthon of dehydropeptide unit.

Recently, much attention has been focused on the synthesis of dehydropeptide (DHP), containing one or more α -dehydroamino acid (DHA) residues, and the correlation between the structure and the bioactivity of DHP.¹⁻⁵⁾

So far, we have developed a few synthetic methods for DHP by the stepwise elongation of DHA with α -amino acid or another DHA⁶⁾ and by the β -elimination of peptide having a leaving group.⁷⁾ In the preceding paper,⁸⁾ we reported briefly the synthesis of N-carboxy α -dehydroamino acid anhydride (Δ NCA)⁹⁾ from N-benzyl-oxycarbonyl-DHA and the application of Δ NCA to the dehydrooligopeptide synthesis without using any coupling reagent. Here, we will report the very useful synthetic method for a variety of DHP by using N-acyl Δ NCA as synthon of DHA or DHP unit.

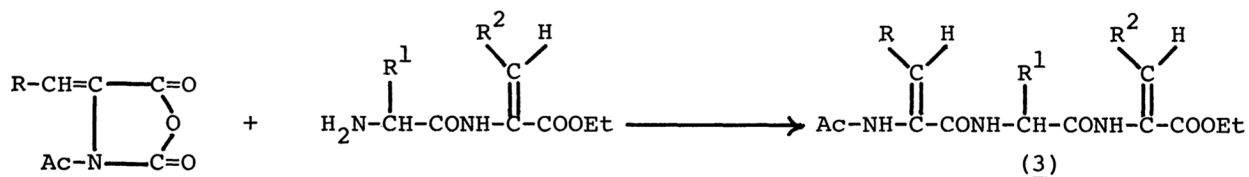
A solution of N-acetyl (Z)- Δ NCA (1), prepared by the acetylation of (Z)-geometric Δ NCA (0.3 mol) with acetyl chloride (0.4 mol) in THF (40 ml) by the usual way,⁸⁾ was treated with (L,L)-dipeptide methyl ester (0.3 mol) in THF (20 ml) at 0-10 °C for 1 hr. The resulting solution was made basic to pH 8-9 with triethylamine. After removal of solvent, the residue was dissolved in ethyl acetate (100 ml) and washed successively with 3% HCl, saturated NaHCO₃ aqueous solution, water, and then dried over anhydrous Na₂SO₄. The evaporation of ethyl acetate gave a crude solid residue, which was purified on a silica gel column using a mixture of benzene-acetone (10 : 1 v/v) as the eluent to give (Z,L,L)-dehydrotripeptide (2) as colorless needles.



Ac- Δ Phe-Leu-Val-OMe	82%	193-194 °C	$[\alpha]_D$	+16.2°
Ac- Δ But-Phe-Val-OMe	95%	135-136 °C	$[\alpha]_D$	-38.3°
Ac- Δ Phe-Phe-Val-OMe	95%	158-159 °C	$[\alpha]_D$	-64.2°
Ac- Δ Leu-Ala-Val-OMe	80%	180-181 °C	$[\alpha]_D$	-22.2°

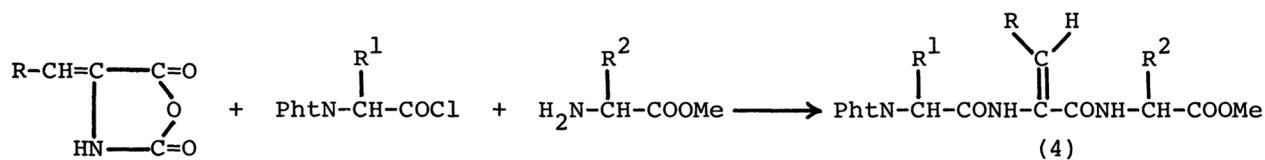
Treatment of 1 with (L,Z)-dehydrodipeptide ethyl ester⁶⁾ followed by similar

work-up gave (Z,L,Z)-dehydrotripeptide (3) as colorless needles.



Ac-ΔLeu-Phe-ΔLeu-OEt	60%	109-110 °C	[α] _D	-45.9°
Ac-ΔLeu-Ala-ΔBut-OEt	71%	syrup	[α] _D	-11.5°
Ac-ΔPhe-Leu-ΔVal-OEt	66%	198-200 °C	[α] _D	+28.3°

On the other hand, to synthesize the tripeptide containing a DHA residue at center, ΔNCA was acylated with an equimolar phthalyl-(L)-amino acid chloride (0.3 mol) in the presence of triethylamine in THF (40 ml) at 5-10 °C for 1 hr, followed by the coupling with (L)-amino acid methyl ester at room temperature for 1.5 hr. After removal of solvent, the residue obtained was submitted to similar work-up to give (L,Z,L)-dehydrotripeptide (4) as colorless needles.



Pht-Leu-ΔBut-Phe-OMe	71%	151-152 °C	[α] _D	-4.5°
Pht-Phe-ΔBut-Gly-OMe	77%	100-101 °C	[α] _D	-113.4°
Pht-Gly-ΔLeu-Phe-OMe	78%	193-194 °C	[α] _D	-20.2°

In conclusion, it was found that the ΔNCA method developed newly by us was very versatile and applicable to the various dehydropeptide syntheses by combination of α-amino acid and DHA. Further works including the analogous study are now in progress.

References

Specific rotation was measured in methanol (c 1.0) at 25 °C.

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- 9) In this paper, the symbol Δ indicates a double bond of DHA residue.

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