Direct Observation of an Alkoxycarbonylamino Acid O-Acylisourea

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Summary The reaction of equivalent amounts of benzyloxycarbonyl-L-valine and NN'-di-isopropylcarbodi-imide in deuteriochloroform at room temperature is very fast and clean giving an O-acylisourea adduct which is stable

in solution for many hours and which has been characterised by n.m.r. spectroscopy.

It seems to be generally accepted that a highly reactive O-acylisourea is formed during the activation of N-protected amino-acids for peptide synthesis by means of carbodi-imides, and that if this is not consumed promptly by added nucleophile it isomerises to the corresponding unreactive N-acylirea or reacts with further N-protected amino-acid to give symmetrical anhydride and urea. As far as we are aware, there have been no examples of the direct observation of O-acylisoureas, although the indirect evidence for their intermediacy in peptide coupling reactions is considerable 1 . In fact, such adducts are stable for many hours in deuteriochloroform at room temperature. Thus addition of one equivalent of NN'-di-isopropylcarbodi-imide to a 0.15 M solution of benzyloxycarbonyl-L-valine

pose, mainly to the N-acylurea, but this had only proceeded to an extent of ca 60% after 6 days. Attempts to isolate the O-acylisourea, however, led to decomposition. When the same experiment was performed with perdeuterio-

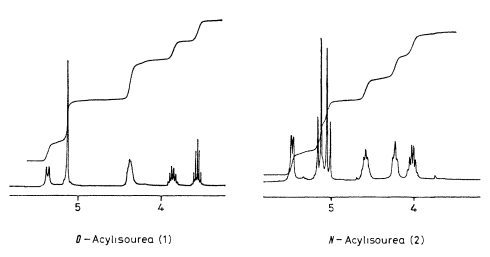


Figure Comparison of the n m r spectra of (1) and (2) in the region δ 3 5—5 5

in deuteriochloroform at room temperature led to rapid clean and complete reaction with formation of O-(benzyloxycarbonyl-L-valyl)-NN'-di-isopropylisourea (1) which was characterised by its 300 MHz n m r spectrum† (see the Figure). The possibility that the spectrum observed was that of the isomeric unreactive N-acylurea (2) was eliminated by its separate preparation and characterisation, its spectrum‡ (see the Figure) was quite different. The spectrum of the O-acylisourea remained completely unchanged for 9 h, on standing overnight it began to decom-

dimethylformamide as solvent sharply contrasting results were obtained. In this case there was no immediate reaction. Even after 30 min both reactants were more than 50% unchanged. A complex mixture comprising mainly N-acylurea (2) NN'-di-isopropylurea and benzyloxycarbonyl-L-valine anhydride was slowly formed, at no stage were peaks attributable to the O-acylisourea (1) detected

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¹ For a review see D H Rich and J Singh in The Peptides ed E Gross and J Meienhofer Academic Press New York, 1979 vol 1, ch 5

† The 300 MHz n m r spectrum of a 0 15 m solution of (1) in CDCl₂ at 20 °C 8 7 35 [s 5H ArH], 5 35 [d collapsing to s on irradiation of the band at 4 2—4 5 1H PhCH₂OCONH] 5 12 [s 2H PhCH₂] 4 2—4 5 [complex 2H (CH₃)₂CHNH and NHCHCO] 3 87 [complex simplifying to a septet on irradiation of the band at 4 2—4 5 1H (CH₃)₂CHNH] 3 58 [septet 1H (CH₃)₂CHN=] 2 15—2 35 [complex 1H (CH₃)₂CHCH] 1 25 and 1 14 [2 × d 12H 2 × (CH₃)₂CHN] and 1 05 and 0 95 [2 × d 6H, (CH₃)₂CHCH] The 0 15 M solution was prepared by addition of 1 equiv of di isopropylcarbodi-imide (the spectrum of which consists in CDCl₃ of a septet at 3 57 and a doublet at 1 23 of relative intensity 1 6) as a neat liquid to a 0 15 M solution of benzyloxycarbonyl L valine in CDCl₃

† The 300 MHz n m r spectrum of a 0 15 M solution of (2) in CDCl₃ at 20 °C δ 7 5 [br 1H NHCH(CH₃)₂] 7 35 [s 5H ArH] 5 45 [d 1H PhCH₂OCONH] 5 07 [ABq 2H PhCH₂] 4 57 [complex 1H NHCHCO] 4 22 [complex 1H (CH₃)₂CHNCONH] 4 02 [complex 1H (CH₃)₂CHNH] 1 93 [complex 1H (CH₃)₂CHCH] 1 36 and 1 27 [2 × d 6H (CH₃)₂CHCH] and 1 24 and 0 98 [2 × d 12H 2 × (CH₃)₂CHN] Compound (2) was isolated chromatographically from the mixture of neutral products resulting from an attempted coupling of equivalent amounts of benzyloxycarbonyl L valine and glycine ethyl ester hydrochloride in dimethylformamide by 1 equiv of di-isopropylcarbodi-imide in the presence of 1 equiv of triethylamine at 0 °C It was obtained as a semisolid material which was stable indefinitely and was homogeneous by t1c and gave 1r ms and analytical data in full accord with the structure (2)