Benzotriazole-assisted Synthesis of Monoacyl α -Aminoglycines

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Adducts (8,12), derived from benzotriazole (7), glyoxylic ester (6) or acid (11), and an amide (5), react with ammonia in a novel, convenient route to monoacylated α -aminoglycines (9,13) useful for the synthesis of peptide analogues.

Reversing one or more of the amide groups (*i.e.* –CHRCONH– to –CHRNHCO–) of a linear peptide gives a so called 'partially modified retro isomer' and represents an important strategy in peptide analogue research. ^{1a,2} The modified sequence requires both a malonic unit and an (much less easily available) α, α -diamino moiety. Such α, α -diamino units have been synthesized by Curtius-^{1,3,5} or Hoffmann-type^{2,3,4} rearrangement of appropriately protected amino acid

derivatives. Recently, Bock and co-workers⁶ obtained protected α -aminoglycines (4) from α -hydroxy-N-(benzyloxycarbonyl)glycine (1) via intermediates (2) and (3) (Scheme 1); direct reaction of (2) with NH₃ did not give (4).

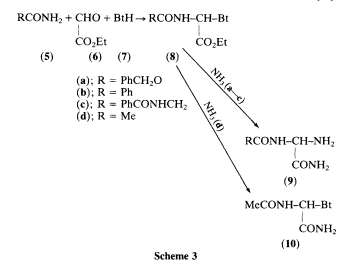
Earlier we reported⁷ convenient syntheses of compounds of type >N-CHR-X mediated by benzotriazole by the general route of Scheme 2. More recently, this methodology with ethyl glyoxylate as oxo-component ($R = CO_2Et$) and organozinc

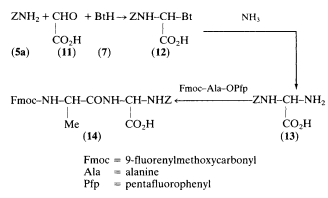
Scheme 2

carbanion nucleophiles (Nu = alkyl, aryl) allowed the synthesis of α -dialkylaminoesters. Extension of this work to N-nucleophiles has now led to the first direct synthesis of monoacyl- α -aminoglycines.

Condensations of various amides (5a-d), benzotriazole (7) and ethyl glyoxylate (6) (in toluene and p-toluenesulphonic acid catalyst) under Dean-Stark conditions gave adducts (8a-d) in 70-75% yield (Scheme 3). Amide (5a), benzotriazole and glyoxylic acid (11) (in benzene) similarly gave (12) (78%, Scheme 4). Reactions of electrophilic acylglycine synthons (8b,c) and (12) with methanolic NH₃ afforded the α -amino-N-acylglycines (9b,c) and (13) in good yields (86, 98) and 70% respectively), simultaneous amidation of the ester function having occurred. Similar reaction of (8a) afforded (9a) together with dimer product; in aqueous ammonia (9a) was isolated in 37% yield. The work-up procedure is easy: simple filtration in the aqueous case, or evaporation of the methanol and isolation with an appropriate solvent [acetone for (13), ether for the others], afforded solid products in a practically pure state. The side-product benzotriazole was removed in the mother liquor. The reactivity of the α -carbon centre depends on the character of the acyl group (RCO): in the case of R = Me(8d), overnight reaction with methanolic NH₃ at 5 °C led almost selectively to amide (10). However, some formation of benzotriazole was observed (TLC) in these reactions as well, indicating that the replacement of the Bt-moiety should be attainable under the appropriate conditions. Compounds (9a—c) and (13) are stable at room temperature, however, attempted recrystallization of (9a) from boiling EtOH led to quantitative dimerization to [ZNHCH(CONH2)]2NH.

Application of the synthesized α -amino-N-acylglycines for peptide synthesis was tested by reaction of amino acid (13) with fluorenylmethoxycarbonylalanine pentafluorophenyl ester (Fmoc–Ala–OPFP) under the usual conditions to yield the expected dipeptide Fmoc–Ala–Gly(NHZ)–OH (14). However, the preparation of such peptides possessing the α -aminoglycine unit in the C-terminal position is more conveniently effected directly by our benzotriazole-assisted glycination method as illustrated by the synthesis of dipeptide (9c).†





Scheme 4

Preliminary results indicate that this method can be applied to primary and secondary amines as nucleophiles, as well as to simple aliphatic and aromatic aldehydes as oxo-components. These extensions should provide a general route to monoacyl- α, α -diamino compounds.

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 $[\]dagger$ All new compounds were fully characterized by C,H,N analyses and by $^{13}\text{C-}$ and $^{1}\text{H-n.m.r.}$ spectra.