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α -Methylphenacyl thioesters as convenient thioacid precursors†

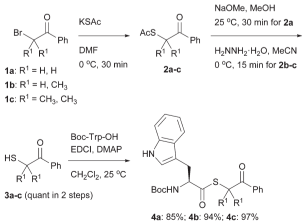
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α-Methylphenacyl (Mpa) thioesters are described as precursors of thioacids. Mpa thioesters are accessible via the condensation of carboxylic acids and phenacyl thiol, which is easily prepared without column chromatography. The Mpa thioesters are selectively deprotected by reduction with zinc dust in the presence of conventional thioacid protecting groups. In addition, the Mpa group exhibits orthogonal reactivity to the Boc group. These features are expected to facilitate the preparation of complex thioacids, including those in peptides.

The thioacid functional group, due to its unique reactivity, can be an expedient handle by which to convergently and chemoselectively install modification modules, particularly in peptides. For example, thioacid-based amide ligation reactions relying on unprotected aziridines,2 electron-deficient azides,3 electron-deficient arylsulfonamides,4 isonitriles,5 and isocyanates⁶ have been reported. Furthermore, condensation reactions between thioacids and amines have also been described recently, employing the Sanger and Mukaiyama reagents,⁷ HOBt, organonitrite, Cu(II), HOBt-Cu(OAc)2, and N,O-bis (trimethylsilyl)acetamide as activators.12

A promising approach to preparing peptide thioacids is to carry thioesters through the peptide elongation sequence and then release the thioacids by reliable and simple reactions. Currently, several thioesters are available as thioacid precursors, including 2,4,6-trimethoxybenzyl (Tmob),3 9-fluorenylmethyl (Fm), 13 trityl (Trt), 14 and 2-cyanoethyl, 15 which release thioacids under either basic or acidic conditions. As for C-terminal peptide thioacids, other approaches were reported, which utilize N-S acyl transfer16-18 or acyl hydrazines.19 We were interested in a different type of precursor that would permit a wider range of applications for thioacid chemistry,

and herein, report a novel thioacid precursor that has, notably, reactivity orthogonal to the Boc group. As thioesters are not generally stable to nucleophilic amines such as piperidine, Boc-based elongation would be the first choice for preparing thioacid-bearing peptides. Therefore, thioacid precursors with reactivity orthogonal to the Boc group could be useful for the preparation of peptide thioacids. In this context, we focused on phenacyl groups. As a protective group for carboxylic acids, phenacyl esters have been used due to their stability under acidic conditions and removal by one-electron reduction in mildly acidic media.²⁰ However, phenacyl esters are known to be relatively electrophilic and susceptible to undesired transesterification. Therefore, we prepared phenacyl thiols 3a-c, which vary in bulkiness (Scheme 1), to examine the stabilities of their thioesters. Each variant was prepared from the commercially available phenacyl bromides 1a-c by a two-step sequence of substitution with thioacetate to give 2a-c, followed by deacetylation. These are all easily scalable spot-to-spot reactions, and afford phenacylthiol 3a, α-methylphenacylthiol 3b, and α,α-dimethylphenacylthiol 3c quantitatively, without the need for column chromatography.



Scheme 1 Preparation of thiols 3a-c and thioesters 4a-c

DMF 0 °C, 30 min 1a: R1 = H, H 1b: R1 = H. CH2 1c: R1 = CH3, CH3 Boc-Trp-OH

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Each thiol was efficiently coupled with Boc-protected tryptophan using standard coupling procedures in the presence of DMAP.²¹ The lower yield in the preparation of 4a is due to the partial loss of 3a observed during silica gel column chromatography. Thioesters 4a-c were subjected to reducing conditions using zinc dust in aqueous acetic acid as a benchmark (Table 1). The efficiency in liberating the thioacid was evaluated after conversion to the corresponding benzyl thioester 5, because thioacids generally have two tautomeric forms (O- and S-acids) that complicate spectroscopic analysis. 22 All three substrates were deprotected to give the corresponding thioacid. The reaction rate clearly depended on the bulkiness of the phenacyl moiety. The bulkier thioester 4c required a longer reaction time than 4a and 4b, even though the intermediate phenacyl radical of 4c would presumably be more stable than those of the others. Elevating the reaction temperature shortened the reaction time and improved the yield. We considered the α-methylphenacyl group, abbreviated as Mpa, to be a good entry as a precursor for thioacids.²³

Table 2 summarizes the scope of this thioacid preparation procedure. Each of the aromatic, aliphatic, and amino acids 6a-i was converted to the Mpa thioester 7a-i using EDCI and DMAP in excellent isolated yields. These precursors afforded the corresponding thioacids smoothly when treated with zinc at ambient temperature. Excess zinc dust was removed by filtration through a silica gel pad, and then the solvent was evaporated to effectively give the thioacid, according to the yield after benzylation. It is noteworthy that no diastereomeric benzyl thioester of 8f was observed, which excludes the possibility of racemization in the reaction sequence. The present transformation was also applicable to Fmoc-protected amino acid 6g. Furthermore, carboxylic acids having additional thioester groups such as 6h and 6i could be converted to the bisthioesters, and the Mpa group could be removed in the presence of either an Fm- or Tmob-thioester. The thioacids could also be captured by 3-bromopropionitrile instead of benzyl bromide to give the 2-cyanoethyl thioester 8i'.

Table 1 Deprotection of phenacyl thioesters 4a-c

Entry	Thioesters 4a–c	Temp./°C	Time	Yield of 5 ^{<i>a</i>} /%
1	4a	25	1 h	92
2	4a	40	20 min	88
3	4b	25	3 h	86
4	4b	40	1 h	86
5	4c	25	26 h	50
6	4c	40	6 h	83

a Isolated yields.

Table 2 Substrate scope of thioacid preparation via Mpa thioesters

		1)) Zn	
	3b (= <i>Mpa</i> SH)		90% AcOH aq.	
	EDCI, DMAP	Ира	25 °C	
R ² -COOH	─			R ² -COSBn
6a-i	CH ₂ Cl ₂ , 25 °C O	7a-i ²⁾) BnBr, Cs ₂ CO ₃ DMF	8a-i

Entry	R ² -COOH	Yield of 7a–i ^{<i>a</i>} /%	Yield of 8a-i ^a /%
1	PhCOOH (6a)	90	80
2	ⁿ C ₉ H ₁₉ COOH (6b)	92	91
3	Boc-Phe-OH (6c)	94	83
4	Boc-Gly-OH (6d)	95	83
5	Boc-Ser(OBn)-OH (6e)	98	78
6	Boc-Thr(OBn)-OH (6f)	94	73
7	Fmoc-Trp-OH (6g)	98	85
8	Boc-Glu(SFm)-OH (6h)	74	78
9	Fmoc-Glu(STmob)-OH (6i)	77	81
10	6i	_	77^{b}

^a Isolated yields. ^b 3-Bromopropionitrile was used instead of BnBr, which afforded 2-cyanoethyl thioester 8i'.

Next, we examined the orthogonality of the Mpa thioesters and Boc groups. As shown in Scheme 2, Boc-protected thioglycinate 7d was employed as the C-terminus, and peptide elongation based on Boc-TFA chemistry was conducted. In the

NHBoc

Scheme 2 Synthesis of oligopeptide α -thioacid

presence of the Mpa thioester, the Boc group was successfully deprotected in 40% TFA/CH₂Cl₂. The subsequent coupling reaction afforded dipeptidyl thioester 9, where NMM performed better than DIPEA as a base. Another deprotectioncoupling sequence was performed with lysine having a Boc group at the ω-amine to give tripeptide 10 without any problems. In the presence of the ω-Boc group, Mpa was cleanly removed using zinc at 40 °C to afford the peptidyl thioacid in good yield, which was confirmed by derivatization to the benzyl thioester 11.

Finally, as shown in Scheme 3, we prepared ω-Mpa thioglutamate 16 from the known glutamate 12 by protecting group manipulation, i.e., allylation of the α-carboxylic acid, deprotection of the ω-9-fluorenylmethyl ester, thioesterification using MpaSH, and deallylation using a Pd catalyst. This glutamate residue could be incorporated into a peptide by Boc-TFA chemistry without loss of the Mpa group to afford tripeptide 18. Again, the Mpa group could be removed in the presence of

.OFm AllBr, K2CO3 Piperidine/DMF DMF, 25 °C 25 °C, 20 min BocHN **BocHN** 6 h, 97% 85% ö 12 13 SMpa MpaSH, EDCI Pd(OAc)₂ DMAP PhSiH₃, PPh₃ CH2Cl2, 25 °C CH2Cl2, 25 °C **BocHN** BocHN 2 h, 80% 1 h, 98% 14 15 .SMpa Gly-OEt·HCl SMpa EDCI, HOBt·H2O NMM CH₂Cl₂ BocHN 25 °C, 3 h, 82% 16 SMpa 1) TFA, CH₂Cl₂ **BocHN** 25 °C, 10 min 2) Boc-Trp-OH EDCI, HOBt·H₂O NMM, CH₂Cl₂/DMF 18 25 °C, 2 h, 72% 1) Zn 90% AcOH aq 40 °C, 6 h 2) BnBr, Cs2CO3, DMF 25 °C. 79% in 2 steps 19

Scheme 3 Synthesis of oligopeptide having thioacid on its side-chain.

the Boc group, and benzyl thioester 19 was obtained after treatment with benzyl bromide. This orthogonality permits the preparation of peptidyl thioacids at either the C-terminus or the side-chain.

Conclusions

α-Methylphenacyl (Mpa) thioesters were developed as precursors for thioacids. MpaSH is easily accessible from commercially available, low-cost materials and readily introduced to carboxylic acids. Mpa thioesters can facilitate the preparation of peptides bearing a thioacid either at the C-terminus or on the side chain. The orthogonal reactivity to the Boc-group and chemoselectivity over Tmob- and Fm-thioesters are expected to enable the preparation of complex thioacids such as peptide thioacids. We are now applying this chemistry to the preparation of oligo- and polypeptides.

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- 23 In the reaction of **4b**, conversion to the corresponding thioacid was verified by ESI-HRMS; calcd for $C_{16}H_{19}N_2O_3S^ [M-H]^-$ 319.1122, found 319.1103.