

Chemistry as an Expanding Resource in Protein Science: Fully Synthetic and Fully Active Human Parathyroid Hormone-Related Protein (1–141)**

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Human parathyroid hormone-related protein (hPTHrP), originally isolated from lung cancer cell lines in 1987,^[1] is a 141-amino acid polypeptide widely found in both normal and tumor tissue cells. The N-terminal region of hPTHrP possesses a high degree of structural homology with human parathyroid hormone (hPTH), and both hormones effect the elevation of calcium levels in the blood.^[2] Although PTH and PTHrP act through binding to the same receptor, the PTH-receptor type-1 (PTHr1), *in vitro* studies suggest that the two ligands may differ in the precise molecular modes of their receptor interactions.^[3,4] Under normal conditions, hPTHrP, which is widely expressed in the tissues of embryos and adults, plays an essential role in a range of functions related to development and growth, including fostering of the cartilaginous growth plate,^[5] bone anabolism,^[6] development of mammary gland,^[7] transport of calcium ions across the placenta,^[8] relaxation of smooth muscle, or vasodilatation,^[9] and eruption of tooth.^[10] In analogy to the related anti-osteoporosis therapeutic agent PTH, researchers have found that PTHrP, when administered daily, may induce anabolic effects on the skeleton. Interestingly, the risk of hypercalcemia associated with PTH-based therapeutics may be lowered with the use of PTHrP. These findings raise the possibility that PTHrP and/or congeners thereof could offer an advantage over currently used PTH peptides in therapeutics related to osteoporosis.

A growing understanding of the role that hPTHrP may play in mediating the progression of cancer further enhances interest in this polypeptide. An intriguing property of

hPTHrP is the finding that it exhibits anti-apoptotic and proliferation-promoting effects on tumor cells.^[11–14] Recent studies have shown that antagonists of PTHr1 are able to remarkably inhibit the growth of tumors.^[15–17]

The development of an efficient synthetic route to homogeneous hPTHrP, and analogues thereof, would facilitate the systematic study of the interaction between hPTHrP and its receptor PTHr1. Such research would offer important insights into the structure–activity relationship (SAR) of the polypeptide, and could well facilitate the development of practical PTHr1 antagonists to suppress the growth of tumors, or agonists for the treatment of osteoporosis.^[18] Certainly, one could imagine that a wisely crafted hPTHrP lookalike could have exploitable antiproliferative properties.

In our judgment, the synthesis of protein targets offers significant learning opportunities at the interface of chemistry, biology, and medicine.^[19] The advantage of pursuing chemistry-based approaches to protein targets arises from the fact that this forum uniquely allows the versatile design of unnatural probe structures possessing defined alterations of amino sequence and structure, including the incorporation of nonproteogenic amino acids.^[20–22] Notwithstanding impressive accomplishments in protein engineering, which were enabled by spectacular advances in molecular biology, we have felt that chemical-based synthesis, in principle, also has much to offer in terms of reaching a specific protein target, in reasonable research-level quantities (usually several milligrams), above all with very high levels of homogeneity. Thus, the purposes of this research were several. First, we hoped to reach hPTHrP by purely chemical means, and to show that it manifests full biological function. With this task accomplished, the basis for an SAR program, involving alterations of primary structure (proteogenic and nonproteogenic amino acid substitutions) and molecular constraints, would be in place. More broadly, we would be exploring, albeit in only a preliminary fashion, prospects for using chemistry as a major resource in protein discovery.^[23]

The field of protein chemical synthesis was greatly advanced with the discovery of cysteine-based native chemical ligation (NCL) by Kent and co-workers.^[24–26] More recently, the scope of NCL has been expanded to encompass a wide range of noncysteine amino acids, through methods developed in our laboratory and others.^[27–33] The general noncysteine-based NCL strategy adopted by our group involves the installation of a temporary thiol functionality on the N-terminal amino acid residue at the site of ligation

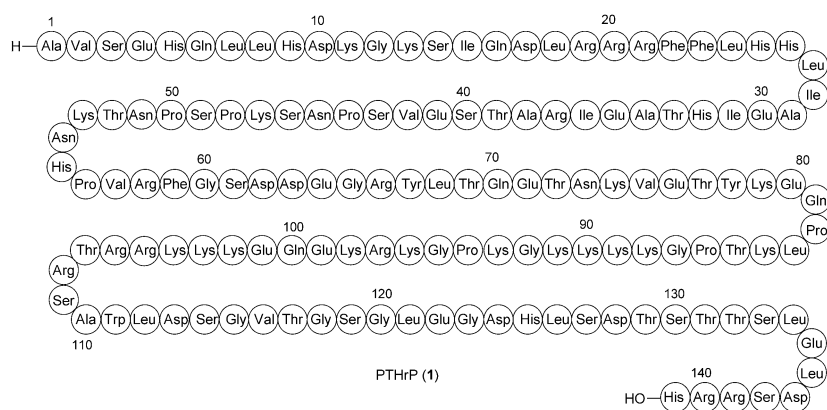
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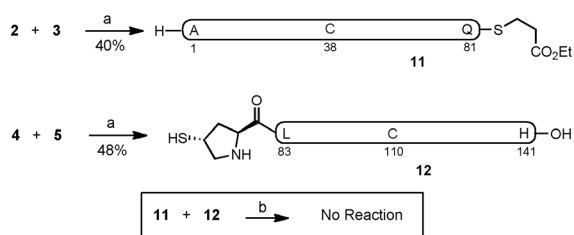


Figure 4. a) 6 M Gn-HCl, 100 mM Na₂HPO₄, 50 mM TCEP, pH 7.2; b) 6 M Gn-HCl, 300 mM Na₂HPO₄, 200 mM MPAA, 20 mM TCEP, pH 7.2. Gn = guanidine, MPAA = 4-mercaptophenylacetic acid, TCEP = tris(2-carboxyethyl)phosphine hydrochloride.

(Figure 4). Similarly, peptides **2** and **3** were ligated under NCL conditions to afford the desired peptide **11** in 40% yield.

We next sought to merge fragments **11** and **12**, en route to hPTHrP, through proline-based ligation.^[40] However, in the presence of 4-mercaptophenylacetic acid (MPAA) catalyst under our standard reaction conditions,^[41] no product was observed after 7 h at room temperature. This result was somewhat surprising, given that the proposed ligation pattern, that is, C-terminal Gln and N-terminal Pro, had been previously demonstrated in our laboratory, albeit in somewhat smaller peptide substrates.^[40] Clearly, our initial synthetic plan toward hPTHrP would require reconfiguration.

In reexamining the primary structure corresponding to hPTHrP, we elected to shift the site of the final ligation from Gln⁸¹–Pro⁸² to Tyr⁶⁷–Leu⁶⁸. We anticipated that application of our recently described formal leucine-based ligation protocol would enable this proposal.^[42] This shift in the disconnection site would require the assembly of fragments **13** and **14**, in place of peptides **3** and **4** (Figure 5). In fact, this alternative disconnection strategy proved quite advantageous from the standpoint of peptide synthesis. Thus, while the synthesis of the previous precursor fragment **3** had required the coupling of two shorter peptides, the new substrates **13** and **14** were both readily accessed through direct SPPS with high purity. Notably, although peptide **14** is quite long, the lysine-rich regions of this segment serve to alleviate potential issues of peptide aggregation.

The fully protected peptides **15** and **16** were synthesized by Fmoc-based solid-phase peptide synthesis (0.1 mmol scale, Figure 6). The preleucine surrogate was incorporated onto

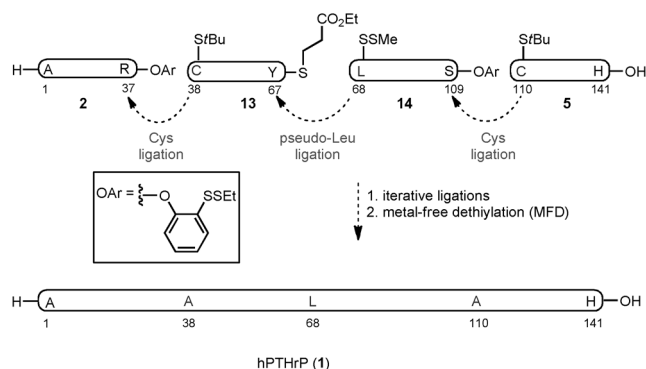


Figure 5. Alternative retrosynthetic plan toward hPTHrP (1-141)

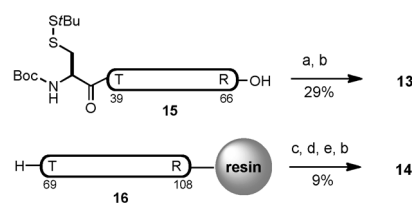


Figure 6. a) H-Tyr(*t*Bu)-S-CH₂CH₂CO₂Et, EDC, HOObt, CHCl₃/TFE = 3:1 (v/v); b) TFA/PhOH/*i*Pr₃SiH/H₂O = 88:2:6:4 (v/v); c) (3*S*)-N-Boc-3-CH₂SS-Leu-OH, HATU, *i*Pr₂EtN, DMF; d) HOAc/TFE/DCM = 1:1:8 (v/v); e) H-Ser(*t*Bu)-O-(2-EtSS)Ph-HCl, EDC, HOObt, CHCl₃/TFE = 3:1 (v/v). Boc = *t*-butoxycarbonyl, DMF = N, N-dimethylformamide, HATU = 1-[bis(dimethylamino)methylene]-1*H*-1,2,3-triazolo[4,5-*b*]pyridinium 3-oxide hexafluorophosphate. Peptide fragments **15** and **16** (in bold) contain protected amino acid residues.

the N-terminus of the fully protected peptide through application of the previously described method. Finally, amino acid residues presenting C-terminal thioester moieties for NCL were attached to the peptides through recourse to standard protocols.

In the event, NCL of peptide segments **2** and **13** was conducted in a pH 7.2 buffer at room temperature, to give the desired peptide **17** in 56% yield after 3 h (Figure 7). Segments

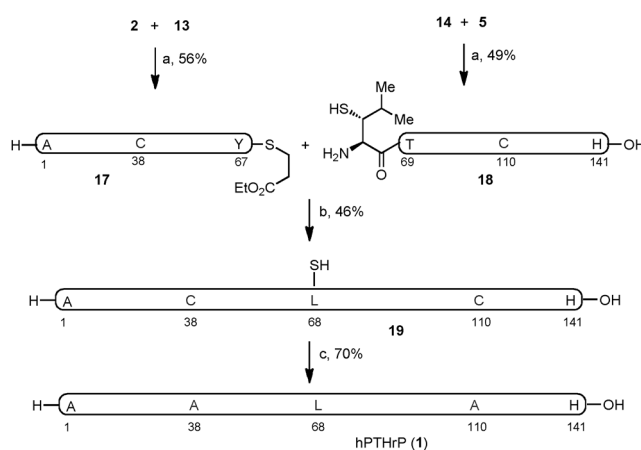


Figure 7. a) 6 M Gn-HCl, 100 mM Na₂HPO₄, 50 mM TCEP, pH 7.2; b) 6 M Gn-HCl, 300 mM Na₂HPO₄, 200 mM MPAA, 20 mM TCEP, pH 7.2; c) TCEP, *t*BuSH, VA-044, 37°C. VA-044 = 2,2'-azobis[2-(2-imidazolin-2-yl)propane]dihydrochloride.

14 and **5** were similarly ligated to generate **18** in 49% isolated yield. In the key coupling event, peptide segments **17** and **18** smoothly underwent the hoped-for thio-Leu ligation in the presence of 4-mercaptophenylacetic acid (MPAA) in pH 7.2 buffer to yield peptide **19** in 46% isolated yield after 20 h. Finally, under the metal-free radical desulfurization conditions developed in our laboratory (VA-044, TCEP pH 7.2 buffer at 37°C), peptide **19**, which bears three extraneous thiol groups, was readily converted to the target compound PTHrP(1-141) in 70% isolated yield (Figure 7 and Supporting Information). Under native conditions, the synthetic hPTHrP(1-141) folded spontaneously,^[43,44] as demonstrated

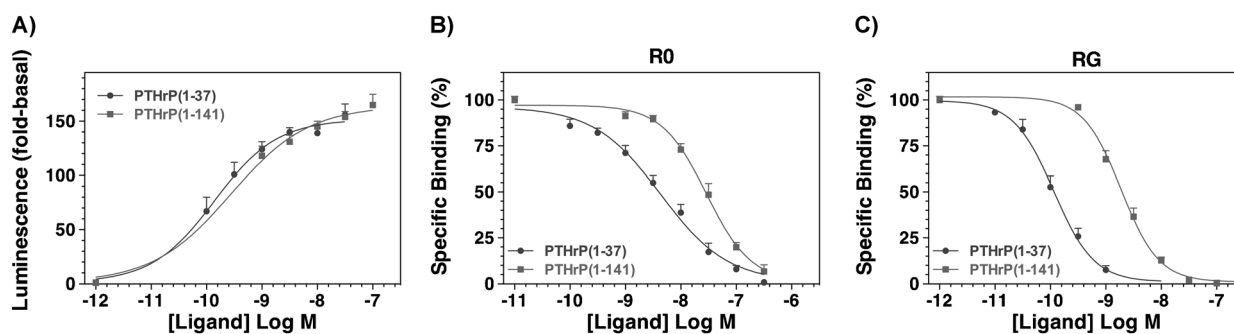


Figure 8. The biological properties of hPTHrP(1-37) and hPTHrP(1-141) assessed in vitro.

Table 1: Functional properties of intact and N-terminal PTHrP peptide ligands.

Ligand	Binding						cAMP (Glosensor assay)						
	R ⁰		P	RG		n	Emax fold-basal ^[a]	pEC ₅₀					
	pIC ₅₀			pIC ₅₀				pEC ₅₀					
	nm	fold		nm	fold			nm	fold	p	n		
PTHrP(1-37)-OH	8.40 ± 0.12		1.00	9.98 ± 0.10		1.00	4	161 ± 21	1.000	9.68 ± 0.37		1.000	3
	3.98	1.0		0.105	1.0					0.21	1.0		
PTHrP(1-141)-OH	7.53 ± 0.11		0.002	8.66 ± 0.08		0.0001	4	175 ± 20	0.670	9.39 ± 0.39		0.610	3
	29.51	7.4		2.19	21					0.41	2.0		

[a] Basal = 1.223 ± 145 cps, *n* = 3.

in the circular dichroism (CD) measurements (190–250 nm, Supporting Information).

The functional properties of the PTHrP(1–141) polypeptide were evaluated in vitro using cells expressing the hPTHrP1. When assessed in an HEK-293-derived cell line that stably expresses the hPTHrP1 along with the Glosensor cAMP reporter gene construct,^[45] the PTHrP(1–141) peptide induced the formation of cAMP with the same potency (EC₅₀) and efficacy (Emax) as did PTHrP(1–37) (Figure 8A; Table 1).^[46] The affinity with which these ligands bound to the hPTHrP1 was assessed in membrane-based competition assays designed to assess binding to two pharmacologically distinct, high-affinity PTHR conformations: a G protein independent conformation (R⁰) and a G protein dependent conformation (RG).^[47] Assays for R⁰ were performed using ¹²⁵I-PTH(1–34) as a tracer radioligand, and an excess of GTPγS, which uncouples receptor/G protein complexes, was added to the reactions. Assays for RG binding were performed using ¹²⁵I-M-PTH(1–15) tracer radioligand and membranes from cells expressing a high affinity, Gas mutant. Under either of these conditions, the PTHrP(1–141) peptide bound with an affinity that was sufficient to fully compete with the tracer radioligand for binding to the receptor, but nevertheless was moderately weaker than that of PTHrP(1–37) (Figure 8B and C; Table 1). The reasons for this weaker apparent binding of the full-length peptide to the receptor, as compared to the shorter-length N-terminal fragment peptide, despite a similar potency for cAMP formation, is not clear at present. Indeed, crystal-structure analysis of the complex formed between a PTHrP peptide and the N-terminal binding domain of the PTHR1 indicates that most, if not all, of the key binding interactions that occur between the ligand and this region of the receptor involve residues limited to the (20–34) region of the ligand.

In summary, the first total chemical synthesis of hPTHrP(1–141) has been achieved. This highly convergent route features iterative, noncysteine-based native chemical ligations, and metal-free desulfurization. The synthesis was accomplished in a convergent fashion. The total yield was 16% from peptide **14**. This efficient synthetic strategy will now be used as a means by which to produce significant quantities of homogeneous hPTHrP(1–141), and congeners thereof, to facilitate hPTHrP- and PTHR1-directed research in the fields of oncology and osteoporosis therapeutics.

More broadly, it seems to be the case that, from a pharmaceutical protein discovery perspective, chemical synthesis may well provide a faster first-time synthesis of a significantly sized all-proteogenic protein in higher levels of purity than that available through molecular biology means of expression. This is even more the case with significantly sized glycoproteins, let alone proteins that carry unnatural amino acids or specifically designed implements for controlling secondary structure.

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