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Bioorganic & Medicinal Chemistry Letters 16 (2006) 5199–5202

Bioorganic & Medicinal Chemistry Letters

Inhibition of HIV-1 integrase activity by synthetic peptides derived from the HIV-1 HXB2 Pol region of the viral genome

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Received 27 May 2006; accepted 5 July 2006 Available online 1 August 2006

Abstract—Peptides deriving from the HIV-1 HXB2 Pol gene sequence were evaluated for inhibitory activity against wild-type (WT) and mutant HIV-1 integrase (IN). The most potent peptide corresponding to a region on the reverse transcriptase (RT) subunit of the Pol polyprotein showed IC₅₀ value of 5 and 2 μ M for 3'-processing and strand transfer, respectively. These peptides, and their analogs, may potentially be used in the elucidation of structural and functional epitopes of IN involved in protein–protein and protein–small molecule interactions.

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The Pol genomes of retroviruses encode three essential enzymes—protease (PR), reverse transcriptase (RT), and integrase (IN). The integration of reverse-transcribed human immunodeficiency virus (HIV) genome into host cell DNA is an essential step in viral replication. This process is catalyzed by IN in two steps.¹ After viral DNA assembly on IN, two nucleotides adjacent to a conserved CA dinucleotide are cleaved from each 3'-end of double-stranded viral cDNA (3'-processing). In the subsequent strand transfer step, IN nicks the chromosomal DNA in staggered positions and joins the 3'-end of the newly processed viral DNA into the 5'-end of the chromosomal DNA nicks. Host repair enzymes such as human Rad 18 are thought to complete the 5'-end joining reaction to produce the provirus.^{2,3} These catalytic steps occur in the cytoplasm and the nucleus of infected cells, respectively. Integration and events prior to it are believed to occur as part of a pre-integration complex (PIC), in which IN, RT, and other essential viral and cellular proteins interact, in the cytosol (early PIC) and in the nucleus (late PIC).^{4,5} To date, validated viral and cellular co-factors of IN such as RT have been shown to both stimulate integration in vitro and in vivo^{6,7} as well as inhibit

integration.^{4,8–10} The cellular and viral proteins that interact with IN each constitute a promising new therapeutic target in combating the virus.¹¹ Elucidating the structural and catalytic nature of such interactions is thus important, especially given that no crystal structure exists of full-length HIV-1 IN.

There is an interest in the possibility of designing inhibitors of IN that target cellular protein–IN and viral protein–IN interactions. ^{12–14} Target-sequence based disruption of functional protein-protein interactions by IN inhibitory peptides or their derivatives should adversely affect productive viral activity. It has been suggested that conformational analysis of peptides will allow the development of peptidomimetic and other inhibitors against IN and that peptide fragments reproducing critical functional epitopes would disrupt protein-protein interactions and inhibit biological activity of the enzyme.¹³ Data from our group suggest that synthetic peptides deriving from the IN amino acid sequence implicate residues critical to enzyme structure and function.¹⁴ Peptides may also have potential in clinical settings, given the success of Enfuvirtide, an anti-HIV-1 peptide drug derived from the viral glycoprotein involved in host cell entry. Several peptide inhibitors of IN targeting catalytic activity as well as oligomerization, although far from entering actual drug development, have been reported. 12,15–18 These include synthetic peptides 12,19–21 from various regions of the IN enzyme as well as those deriving from natural products. 22,23

Keywords: HIV-1 integrase; Protein-protein interaction; Reverse transcriptase; Peptide; Mutant.

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Recently, small soluble peptides from the viral activator Tat domain were shown to significantly inhibit viral replication.¹⁷

We have previously presented preliminary data from in vitro screening of a library of 20-mer synthetic peptides deriving from the HIV-1 HXB2 Pol genome, against IN catalytic activity.²⁴ Herein we report the evaluation of these peptides for inhibition of both 3'-processing and strand transfer. Of these, the most potent inhibitory peptides derive from the RT region of the Pol genome (Fig. 1), while those peptides deriving from the PR region were inactive. A separate publication from our group¹⁴ details the inhibitory activity of shorter peptides from the IN region, as well as their derivatives. It should be noted that a study published last year produced results significantly different from ours for the same RT peptides.²⁵

The 20-mer peptides tested against purified wild-type IN spanned the 1003 amino acid length of the Pol polyprotein, with 10-mer overlaps between sequential peptides. The most active peptide **56** (IC $_{50}$ values of 5 μ M for 3′ processing and 2.5 μ M for strand transfer) derives from

the RT connector region of the protein. Two other peptides, **34** and **53**, showing IC₅₀ values in the micromolar range also derive from the RT palm and connector regions. All three are in close proximity to the active site of the enzyme (Table 1). Peptides **65** and **68** deriving from the RNase H region also exhibit inhibition of IN catalytic activity in the low micromolar range, although less potent than the previous three peptides. We also evaluated the five potent peptides against the IN (F185K/C280S) soluble mutant. While this mutant has been shown to be slightly more resistant as compared to wild-type to small molecule IN inhibitors, this is the first reported case of peptides showing similar inhibitory behavior (Table 1).

The full set of HIV-1 HXB2 Pol 20-mer peptides (1.0 mg, lyophilized) was obtained from the National Institutes of Health AIDS Research and Reference Reagent Program Catalog (catalog number 4358). The first amino acid of the first peptide is recorded as amino acid 433 in the full-length genomic sequence of the *Pol* polyprotein in GenBank (Accession No. K03455). Serial dilutions of the peptides were performed in deionized water or DMSO. Original peptides and stock solutions

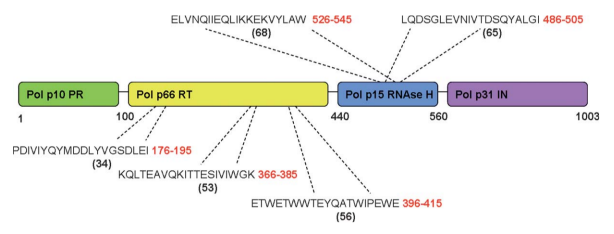


Figure 1. The five most potent peptides derive from different regions of the HIV-1 HXB2 Pol genome. Numbers adjacent to each peptide sequence indicate amino acid spans of the respective subunits on the Pol polypeptide (Numbering according to the HXB2 Numbering Engine nomenclature, Los Alamos National Laboratory). Numbers in parentheses indicate the number of the peptide (domains not drawn to scale).

Table 1. IC₅₀ (μM) values of peptides against wild-type and mutant IN

Peptide	Peptide sequence	Activity	WT^a	SM^a	C130S	C130A
34	PDIVIYQYMDDLYVGSDLEI	3'-Processing	6 ± 1	28 ± 2	41 ± 2	20
		Strand transfer	10 ± 1	23 ± 2	2 ± 1	5
53	KQLTEAVQKITTESIVIWGK	3'-Processing	7 ± 1	51 ± 7	29 ± 1	15
		Strand transfer	4 ± 1	31 ± 7	2 ± 1	10
56	ETWETWWTEYWQATWIPEWE	3'-Processing	6 ± 1	13 ± 2	126 ± 3	25
		Strand transfer	2 ± 1	9 ± 2	27 ± 1	5
65	LQDSGLEVNIVTDSQYALGI	3'-Processing	11 ± 1	>167	18 ± 3	>167
		Strand transfer	2 ± 1	36 ± 14	4 ± 1	20
64	ELVNQIIEQLIKKEKVYLAW	3'-Processing	15 ± 2	113 ± 11	136 ± 5	45
	-	Strand transfer	14 ± 4	83 ± 5	7 ± 1	15

^a Abbreviations: WT, wild-type IN; SM, soluble mutant.

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